

MRIDIUMTM MRI INFUSION SYSTEM

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REF 1124

MRidiumTM 3850 Infusion System Operation Manual, Part Number 1124 Release 7C, 10/2011 PER ECN 000372

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General Information

This document provides the following directions for use:

- Single channel pump that provides a full range of features in a compact, easy to use, linear peristaltic pump.
- Dual channel pump offers the same features while providing two independent infusion pumps in one instrument.
- The Remote Display/Charger Unit allows for remote ability to control the Pump with a total of two channels from outside the MR Scanner.

The system is designed for use in the following patient care areas:

- MRI (0.2 to 3T systems).
- MRI/Recovery.
- Pump operable and safe in up to 1 Tesla (10,000 Gauss) magnetic field.

EMI Statement:

This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to other devices, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures: 1. Reorient or relocate the receiving device. 2. Increase the separation between the equipment. 3. Connect the equipment into an outlet on a circuit different from that which the other devices(s) are connected. 4. Consult the manufacture or field service technician for help.

Mains Disconnection Method:

3850 Pump: Disconnect power cord (1121) from Appliance Inlet on the side of the MRI power supply unit (1120).

3855 Remote/Charger Unit: Disconnect power cord (1128) from Appliance Inlet on the rear of the Unit.

Alternate Voltage/Export:

For power cord plug types see local country distributor. Unit shipped in USA with US 3 pin power NEMA plug.

3850 EQUIPMENT CLASSIFICATION		
Classification according to IEC 60601-1		
According to the type of protection against electrical shock:	Class I equipment and internally powered	
According to the degree of protection against electrical shock:	Type CF (defibrillator-proof) equipment	
According to the type of protection against harmful ingress of water:	Ordinary Equipment. Complies with Section 44.4 of the Infusion Pump standard, IEC 60601-2-24.	
According to the methods of sterilization or disinfection:	Non-sterilizable. Use of liquid surface disinfects only.	
According to the mode of operation	Continuous operation	
Equipment not suitable for use in the presence of flammable anesthetic mixture with air or with oxygen or nitrous oxide		

About the Pump

The MRidiumTM 3850 Infusion System incorporates a unique ultrasonic motor providing the nonmagnetic motor power driving the MRidiumTM 3850 pumping unit. This allows the MRidiumTM 3850 to be constructed with minimal magnetic material and safely operable in high magnetic fields.

Features include

- Non-magnetic ultrasonic drive motor.
- Special aluminium RF noise shielding enclosure.

The pump is equipped with a unique battery display that provides the clinician continuous monitoring of battery capacity available. This information is displayed when the instrument is turned on.

The Dual Rate Feature allows the pump to administer both primary and secondary solutions at separate flow rates and volumes. Using this feature the clinician can select and start a program for secondary delivery. Upon completion of the secondary dose, the pump can automatically switch over to a primary rate. Both channels of the MRidiumTM 3850 can be programmed for primary and secondary operation.

Optional modes are easily accessed with the press of one key.

The Dose Rate Calculator allows the clinician to calculate a dose rate for continuous infusion given concentration and dosage orders.

The Bolus Dose feature allows the clinician to set up an initial infusion rate for a specific Bolus volume, automatically followed by a maintenance rate from the same container.

An optional 2.4 GHz wireless link allows communication between the infusion pump and remote display (denoted by "R" in pump model number).

Qualified service personnel can configure many features of the pump to meet specialized needs.

Specific pump menu screens may vary depending on software release being used.

This device is covered under one or more of the following U.S. Patents:

7,267,661 B2 and International Equivalents

7,404,809 B2 and International Equivalents

7,553,295 and International Equivalents

Other U.S. and International patents pending.

Precautions

Federal Law in the U.S.A. restricts this device to sale by or on the order of a physician.

This device is intended for use by trained medical professionals only.

Refer all service to Iradimed Corporation Authorized Service Representatives.

The 3850 Pump has been specifically designed for operation inside an MRI Magnet Room, and is designed to operate normally in the presence of most frequently encountered electromagnetic interference in the MRI environment. Under extreme levels of interference, such as in close proximity to an electrosurgical generator, cellular telephone or a 2 way radio normal pump operation may be interrupted. Avoid use of this pump under these conditions.

Use only MRI-compatible patient access devices (e.g. needles, luer-ports, etc.) to prevent any possible RF current from reaching the patient's skin.

The Remote Display Charger is intended for use in the MRI control room. Do not operate the 3855 Remote Display/Charger inside the MRI Magnet Room.

For safe operation, use only Iradimed Corporation recommended MRI-compatible or MRI-safe accessories.

The MRI pump must be mounted securely when used in the MRI scan room. Always securely mount the pump using it's integral pole mount clamp or other mechanical fixing means.

Always secure the I.V. Pole wheel locks after positioning within the MRI Magnet Room.

Avoid placing the I.V. Set adjacent to any electrical conductor within the MRI bore, which can become heated during MRI scans.

The Alarm Sound Volume is adjustable for various clinical environments. Ensure the alarm sound level is appropriate for the use environment in the MRI so that it can be heard above the ambient noise level, especially during scanning.

Product damage may occur unless proper care is exercised during unpacking and installation. Do not use the pump if it appears damaged in any way. The battery should be charged before use.

Battery may not be fully charged upon receipt. Connect pump to AC power for at least nine (9) hours before placing the pump into use.

Always connect the infusion pump power supply or Remote Display/Charger to a properly grounded 3-wire power receptacle. If the quality of the earth grounding is in question, use battery power for the infusion pump.

This equipment is not suitable for use in the presence of flammable anesthetic or other gases. Do not use this system in the presences of flammable gases.

Arrange tubing, cords and cables to minimize the risk of patient or other equipment entanglement.

The pump is not intended for high pressure, high viscosity radiological contrast agents.

To avoid patient injury, always respond to Pump or Remote Display/Charger alarms immediately.

Never leave the patient with the Pump stopped if the infusion has not been completed.

To Prevent injury, avoid placement of any NIBP cuff on the limb receiving the I.V. therapy.

The Pump contains materials which must be recycled, or disposed of properly. For proper disposal methods, contact your local sales representative or distributor.

Hospital personnel must ensure the compatibility of the drugs as well as the performance of each pump as part of the overall infusion. Potential hazards such as drug interactions, inaccurate delivery rates, inaccurate pressure alarms and nuisance alarms may arise from other incompatibilities.

Precautions (Continued)

MRI pump uses medical grade silicone rubber and PVC tubing. Do not infuse pharmaceuticals or solutions that are incompatible with these materials (for example, which contain diethylhexylphthalate (DEHP)).

Consult the drug labeling to confirm drug compatibility, concentration, delivery rates and volumes are suitable for concurrent delivery, or piggyback delivery (secondary followed automatically by primary).

Simultaneously infusing with more than one pump into one patient line may significantly affect the infusion rate of at least one of the pumps.

This infusion pump is contraindicated for use on the inlet side of Extra Corporeal Membrane Oxygenation (ECMO) systems or anywhere the negative pressure is greater than -100 mmHg as the high negative pressure can result in uncontrolled fluid flow.

Equipment use outside it's specified environmental conditions can affect infusion accuracy.

Pump Related Precautions

This pump is designed to stop fluid flow under alarm conditions. Periodic patient monitoring must be performed to ensure the infusion is proceeding as expected.

Immediately respond to a "Close Door" alarm due to imminent lapse in infusion therapy.

Immediately respond to the "Check Door" alarm as this indicates possible Free-Flow of IV Fluids.

Dropping, or a severe shock to the pump, could result in damage and resultant pump inaccuracy. Refer the pump to qualified service personnel for proper checkout if either of these conditions occur.

Do not place the pump into use if it fails any of the Power On Self Tests.

Always verify flow rate, VTBI (Volume to be Infused) and/or dose drug entries before starting infusion.

When opening the door, always check that the free flow preventer (black slide clamp) is fully pulled outward to the shut off position.

If questionable Pump operation is observed, or if a System Failure occurs (i.e an unexplained continuous audible alarm with no displayed values), discontinue use of the Pump and refer it to qualified service personnel.

Although unlikely, failure of certain rugged mechanical components, such as the free-flow prevention mechanism could cause fluid delivery limited to the contents of the fluid container. The maximum volume that may be infused under a *single fault condition* is 0.1 mL. Single fault failure of any electronic or motor control component would result in less than 0.3 mL of unexpected fluid delivery.

A small amount of fluid is expelled from the set (less than 0.05 mL) under worse case conditions when the mechanical drive advances to Index each time the pump latch is opened and closed with a set loaded. If potent drugs are being used, take appropriate action to guard against overmedication of the patient.

Always set the occlusion pressure alarm limit at the minimum level required for the prescribed fluid therapy.

Whenever closing the Pump Door, check to make sure that nothing interferes with the pumping mechanism.

During a running infusion, each interruption and re-start of the infusion can add approximately 0.05 mL of delivered fluid to the indicated Volume Infused.

Pump Related Precautions (Continued)

There are dangerous voltages present on internal components that may cause severe shock or death upon contact. Never open the pump casing, it's power supply, or the Remote/Charger when connected to Mains power. Disconnect from AC power and remove battery pack prior to servicing or cleaning.

Do not operate the pump on patients with the battery removed (pump will stop and not alarm during AC power loss without battery installed). Use of a properly maintained and charged battery will allow proper operation. Do not touch battery connector pins and patient simultaneously.

If the Low Battery alarm sounds, connect the pump with its power supply to AC power immediately.

Replace blown fuses on the 1120 Pump MRI power supply or Remote Display/Charger with fuses of the same type and rating only, or a fire hazard could exist.

Never use sharp objects (paper clips, needles, etc.) to clean any part of the pump.

Keep the pump door latch securely closed when the pump is not in use. This will avoid door latch damage.

Do not sterilize the pump or any component by heat, steam, ethylene oxide (ETO), or radiation.

The screen displays the VTBI (Volume to be Infused) in whole integers above 99.9. Any fraction of a milliliter delivered is not displayed, but is retained in memory.

To avoid damage to the I.V. Pump and Pole, always move the I.V. Pole separately from the patient trolley to prevent accidental entanglement.

The pump body is made of aluminum and is non-magnetic. However, when moving the pump within high magnetic fields (>2000 gauss) one might notice Eddy Current effects. These are forces generated in the aluminum which resists motion through the intense magnetic field. Such effects are normal and present no risk of free magnetic movement of the unit.

Set Related Precautions

Always use aseptic techniques. Patient infection could result from mishandled or non-sterile assemblies.

Use only Iradimed Corporation MRidiumTM 1000 Series administration sets. The use of other sets will cause improper pump operation resulting in inaccurate fluid delivery. MRidiumTM 1000 Series administration sets are only intended to be used with the MRidiumTM Pump.

All infusion administration sets are supplied sterile and are single use only. Do not sterilize or reuse.

Prior to use of any Infusion Set, examine the pouch and inspect for damage that could compromise sterility. If the pouch or Set is damaged, discard and use another Set.

Administration sets should be changed per the Center for Disease Control (CDC) guidelines or healthcare provider policy. Discard after use. Design use life of I.V. sets is six (6) hours maximum.

Disconnect the I.V. line from the patient before starting the priming procedure.

Prepare the primary solution container in accordance with the manufacturer's instructions.

The use of positive displacement infusion devices ported together with gravity flow infusion systems into a common I.V. site may impede the flow of the gravity flow system and affect their performance. Hospital personnel must ensure that the performance of the common I.V. site is satisfactory under these conditions.

Set Related Precautions (Continued)

Interconnection with I.V. sets with small inner diameter may affect pump accuracy at high flow rates. Avoid interconnection with small bore diameter I.V. sets (less than 0.050 inch I.V. tubing) if high flow rates are used.

A kinked or occluded I.V. line could cause the pump to operate abnormally and affect the accuracy of the infusion. Before operating this system, verify that the I.V. line is not kinked or occluded.

To avoid nuisance alarms, confirm that the fluid source is positioned higher than the pump.

Setting the primary rate greater than the secondary rate will result in a more rapid infusion of any residual secondary drug remaining in the line, the administration set, and fluid container.

When performing a secondary infusion:

- Secondary solution container must be higher than the primary solution container.
- The Secondary VTBI (Volume to be Infused) setting must be equal to the volume in the secondary container. This requires consideration of such variables as factory overfill, medication additions, etc. Underestimating the volume will cause the remaining secondary solution to be infused at the primary rate; overestimating will result in the primary solution being infused at the secondary rate. Multiple doses from a single container are not possible.

Air bubbles may form distal to the pump as a result of normal out-gassing of dissolved air in the fluid. This may occur if a chilled solution is being used, if the pump is mounted significantly above the patient, or with certain fluids known to routinely outgas. In these cases, an air eliminating filter may be required.

During a prolonged infusion, routinely inspect I.V. Set, access device and patient line assemblies for proper attachment and orientation.

Variations of head height, back pressure, selected catheter type, or any combination of these may affect rate accuracy. Factors that can influence back pressure are: I.V. set configuration, I.V. solution viscosity and I.V. solution temperature. Back pressure may also be affected by catheter type.

The use of pumping infusion devices ported together with gravity flow infusion systems into a common IV site (primary IV set with secondary IV lines) can affect the accuracy of the gravity flow systems, and result in unintended flow rates from these gravity systems. Always ensure the common IV site is acceptable for use under these conditions.

Reference to specific drugs and default parameters are provided for the user's convenience. Always refer to the specific drug product labeling for information concerning appropriate administration techniques and dosages.

Battery Pack Related Precautions

The 1133 Battery Pack contains several lithium-polymer cells and an integral safety circuit. As these cells age, they can expand due to internal gas release, which is anticipated for this type of cell. However, if excessive expansion occurs, this can result in the battery case expanding (swelling), and possibly cause failure of the battery case, cells, or safety circuit. If this is observed, remove the Battery Pack from use and replace it as soon as possible.

The 1133 Battery Pack contains protective circuitry to prevent catastrophic battery failure. If the Battery pack is damaged, this protective circuitry may not prevent battery failure. Remove the Battery Pack from use if the Pack becomes damaged, or the potential for Battery Pack damage is suspected.

Do not use a damaged or swollen 1133 Battery Pack.

Avoid damage to the 1133 Battery Pack by impact, dropping, overheating, or mechanical abuse. Never compress, drop, shock, or strike the 1133 Battery Pack. Never use objects that could puncture the internal battery cells. Any of these actions can cause the battery cells to heat, smoke, or cause catastrophic battery failure, which could result in fire.

Do not attempt to disassemble the 1133 Battery Pack. Damage caused by disassembly or tool use can result in catastrophic battery failure, which could result in fire.

If the 1133 Battery Pack case begins to expand and/or swell, discontinue battery charging and use immediately, and replace the Battery Pack. Continued charging will cause further Battery Pack case expansion, with possible battery case fracture, and potential electrolyte leakage.

If the 1133 Battery Pack becomes damaged, avoid contact with the battery cell electrolyte. If the electrolyte contacts the skin or eyes, seek medical attention immediately.

If the 1133 Battery Pack shows sign of the battery case expanding (swelling), remove the Battery Pack from use and replace it as soon as possible. In extreme conditions, this swelling can cause the 1133 Battery Pack to become jammed or stuck within the 3850 Pump or 3855 Remote Display, and/or cause the Battery Pack plastic case to burst open. If this occurs, do not use tools that could cause damage to the internal battery cells. Refer to the 1125 Service Manual for removal under these conditions.

Under no circumstances should Battery Packs or the internal cells be incinerated as this can cause an explosion.

User Responsibility

This product will perform in conformity with the description contained in this users manual and accompanying labels, inserts, etc., when assembled, operated, maintained and repaired in accordance with the instructions provided. This product must be checked and calibrated periodically. A malfunctioning product should not be used. Parts that are broken, missing, plainly worn, distorted or contaminated should be replaced immediately. Should such repair or replacement become necessary, refer unit to Iradimed Corporation qualified service personnel. This product or any of its parts should not be repaired other than in accordance with written instructions provided by the manufacturer, or altered without written approval of Iradimed Corporation. The user of the product has the sole responsibility for any malfunction which results from improper use, faulty maintenance, improper repair, damage or alteration by anyone other than Iradimed Corporation or Iradimed Corporation authorized service personnel.

Using This Manual

Read this manual completely before attempting to use the pump.

Warning, Cautions and Notes. This manual contains three levels of precautionary information.

- A **Warning** alerts the user that there is a possibility of injury or death to a human being.
- A **Caution** alerts the user that there is a possibility of damage to equipment.
- A **Note** contains essential information deemed especially important by Iradimed Corporation.

Channel A	The designation for the first infusion channel. All pumps contain at least one linear peristaltic pump head for one infusion line.
Channel B	The designation for the second infusion channel. The second infusion pump head is optional and some pumps may not contain it.
hr.	hour.
KVO	Keep-Vein-Open.
mL	milliliter.
Primary	Main Infusant for the prescribed I.V. therapy. The infusion settings that are implemented after any secondary infusion sequence is complete.
Rate	Infusion rate in mL / hr.
Secondary	The first infusion settings to be implemented in an infusion sequence. Sometimes referred to as "piggyback."
VI	Volume Infused in mL.
VTBI	Volume To Be Infused in mL.

Definitions

Symbols



Attention: Consult Accompanying Documents.



Defibrillator Proof Type CF Applied Part



Date of Manufacture



Indicates that the device conforms to the Medical Device Directive



Direct Current



Product is Latex-free



Drops per millimeter specification for I.V. Set is identified on Drop Symbol



Lot or batch code for I.V. Set will be identified near Lot Symbol



Approximate Set priming volume



Main Battery Capacity. (X inside Icon denotes no battery installed.)



AC Power is connected to 100-240 VAC



Secondary Infusion Mode



Volume to be Infused



Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician



Maximum rated load is 20 Kg.



Do not discard. Contact recycler for proper disposal.



Spare Battery Capacity. (X inside Icon denotes no battery installed.)



Appropriate for use in MR environment

MR Conditional. Only appropriate for use in MR environment with manufacturer's defined restrictions.

MR Unsafe. Not appropriate for use in MR environment (i.e. inside the MR magnet room)



Single Use Only



Product Serial Number



Alternating Current



Product does not contain DEHP in the fluid pathway



Product Part Number



Expiration date for I.V set will be identified near hour glass symbol

STERILE R



Contains Lithium. Requires proper disposal/recycling of this material.



Power On, or On



Power Off, or Off



Volume Infused



Input/output connection. Allows data communication.



Input Connection Only Output Connection Only



Storage Temperature Range

deno



EC Authorized Representative



Radio-Frequency transmission source



This product has been certified to UL60601-1 and applicable Particular Standard IEC 60601-2-24, for which the product has been found to comply by Intertek.



2.4 GHz Radio is communicating.



Telecommunications Alert Symbol -Class 2 (European Union Countries only)



This product has been certified to UL60601-1 and applicable Particular Standard IEC 60601-2-24, for which the product has been found to comply by Underwriters Laboratories Inc.

SECTION 1 INTRODUCTION

1.0 Introduction. The MRidiumTM 3850 MRI Infusion System is intended for patients that require medications and/or fluids during an MRI scan. This pump is designed to provide infusion therapy at all stages of the MRI procedure. This system is for use by trained medical personnel only and is not intended for long term patient care outside of an MRI environment.

This system provides the following features:

- Continuous Infusion, Dose Rate Calculation Program and Automatic Bolus.
- Automatic free-flow protection for the I.V. line.
- Up to two channel fluid delivery, each with separate primary/secondary programmable rate and VTBI, into single or separate IV lines.
- Rechargeable long life battery. Lasts up to 12 hours at 125 mL/hr.
- Status indicator light above door (Red for Alarm, Green for Infusing).
- Soft keys to program various functions.
- Large LCD display screen.
- Up and Down arrow control keys to change numeric values quickly and easily.
- Handle for easy portability, weighs less than 11.5 pounds.
- I/O port.
- Optional second channel with the addition of the 3851 SideCarTM.
- Optional remote control with the additional 3855 Remote Display/Charger.
- Memory card slot for easy upgrades.

CAUTION: Handle the MRidiumTM 3850 MRI Infusion Pump and additional channels with care. If it should be dropped or severely jarred, it should be immediately taken out of service and inspected by a qualified biomedical technician.

1.1 Product Description. The MRidiumTM 3850 MRI Infusion System is designed for operation in the MRI environment and may be used on the patient near the MRI magnet (up to the 1.0 Tesla or 10,000 Gauss Line). Operating on battery power, when fully charged this system will provide up to 12 hours of operation at an infusion rate of up to 125 mL/hr and at least four (4) hours at rates up to 999 mL/hr.

The MRidium Wireless Remote Display allows for remote ability to control the Model 3850 MRidium MRI Infusion Pump with a total of two channels from outside the MR Scanner. It utilizes the same user interface as the infusion pumps and will allow adjustment of all pump parameters, rates, titration, volume to be infused, starts, stops, and resetting alarms. The large clear display shows all pump information at your desk top from the control room. The Wireless remote also acts as a charger for a backup or spare battery pack for the 3850 MRidium MRI Infusion Pump. It utilizes a wireless link at 2.4 GHz for easy installation with no image artifacts. The 3850R designates a 3850 Pump with the optional 2.4 GHz wireless link for communication with the 3855 Remote Display.

NOTE: The Remote Display/Charger only operates on AC Mains Power. This unit does not operate on battery power. This unit does not sound a Low Battery alarm for the Spare Battery being charged.

<u>1.1.1</u> <u>Front of the Pump.</u> See Figure 1-1 for the location of the major components on the front of the pump.

- a. **Handle.** Located on the top of the pump, the handle provides for easy transportation of the pump from location to another.
- b. **Main Display.** Provides a visual display of all the parameters and features of the pump.
- c. **Run/Alarm Lamp.** Provides a visual display of the pumps operating condition.
- d. **Battery.** Located behind a door on the back of the pump, the battery provides for power in the absence of a hospital grade AC Mains outlet.
- e. **Pump Area EZ-Latch Door.** Pressing down on this button and lifting this handle unlatches the door protecting the pumping mechanism and infusion line.
- f. Main Control Keypad. Provides control of the various features of the pump.
- g. **Soft keys.** Provides a way to adjust the configuration of the pump.
- h. **AC Power/Battery Charge LED.** Provides indication that MRI power supply is connected and AC power is applied to Charge the pump battery



Figure 1-1. Front of Pump

<u>1.1.2</u> <u>Pump Drive Mechanism.</u> See Figure 1-2 for location of the major components of the infusion system pump drive mechanism.



Figure 1-2. Pump Drive Door Open

- a. **Upper and Lower I.V. Set Snap-In Port.** Provides for a secure mounting of the I.V. set for infusion.
- b. **Linear Peristaltic Pump Drive Mechanism.** Provides for the positive movement of fluids through the infusion line.
- c. **Bubble Detector.** Provides for the detection of air bubbles in the infusion line.
- d. **Free-Flow Detector Switch.** Located internally in the pump's lower snap-in port the IV Set Free-Flow Detector Switch detects when IV set is properly installed and functioning.
- e. **Inlet and Outlet Pressure Sensors.** Provide infusion line pressure measurements for pressure monitoring and alarms.

NOTE: After attachment of the 3851 SidecarTM module (**Refer to Section 4.6**), the operation of Channel B SidecarTM pump drive mechanism is identical to the main pump drive. The only difference is that the Channel B door opens to the left side, which is opposite that of the main pump door.

<u>1.1.3</u> <u>Back of the Pump.</u> See Figure 1-3 (Figure 1-4) for the location of the major components on the back of the pump.

- a. **Handle.** Provides for easy transportation of the pump from location to another.
- b. **Optional Secondary Pump Drive Electrical Connector.** Provides for the addition of a Channel B pump mechanism.
- c. **I.V. Pole Clamp.** Provides a secure mounting of the pump to the I.V. pole.
- d. **Audible Speaker.** Provides the audible sounds for alarms and alerts.
- e. **2.4 GHz Antenna Connector.** Used to connect Antenna for Remote Display/ Charger Option (3850R only).
- f. **Power Input.** Provides for connection to the AC Power Adapter to run the pump on hospital grade AC Mains power. **Connect only model 1120 MRI Power Supply.**



Figure 1-3. Back of Pump (Version 2)

(Serial Number IR5010778 and above)

g. I/O Port. Provides a connection for retrieving the pump's infusion data from memory. Connect only 60601-1 compliant computer for History Log download.

WARNING: Serial I/O Port is not PC-compatible computer format (Pin # 8 is connected to the +5 volt supply). Refer to the Service Manual for History Log data download instructions.

- h. Memory Port. Provides for field update of system software. Use only AM01.
- i. **Battery Compartment.** Provides a safe and secure location for the pump's battery. **Use only Iradimed model 1133 Battery Pack.**

CAUTION: Battery pack is slightly magnetic. Use caution when removing from pump near strong magnetic fields.



Figure 1-4. Back of Pump (Original Version)

<u>1.1.4</u> <u>1120 MRI Power Supply.</u> See Figure 1-5 for information on the pump's MRI Power Supply.

WARNING: AC Adapter is magnetic. Keep outside the 1,000 Gauss line, or at least 10 feet (3 meter) from the MRI magnet. Secure with velcro straps provided to the floor.

NEVER velcro or secure the AC Adapter directly to the pump or I.V. Pole.



Figure 1-5. Model 1120 MRI Power Supply

<u>1.1.5</u> <u>Front of the Remote Display/Charger.</u> See Figure 1-6 for the location of the major components on the front of the 3855 Remote Display/Charger.

- a. **Antenna.** Provides 2.4 GHz bidirectional communication to MRidiumTM 3850 MRI Infusion Pump.
- b. **Main Display.** Provides a visual display of all the parameters and features of the pump.
- c. **Run/Alarm Lamp.** Provides a visual display of the pumps operating condition.
- d. **Battery Charger Compartment.** Located at the top rear of the Remote, allows charging of an 1133 Pump Battery when battery is installed and Remote is connected to AC Mains outlet as indicated by AC Power/Battery Charge LED (located below the power On/Off keys).
- e. Main Control Keypad. Provides control of the various features of the pump.
- f. **Soft keys.** Provides a way to adjust the configuration of the pump.
- g. **AC Power/Battery Charge LED.** Provides indication that AC power is applied (Green) or Charging a spare battery (Amber).
- h. **Memory Port.** Located underneath the front panel provides for field update of system software. **Use only AM01 Software Card**.



Figure 1-6. Front of 3855 Remote Display/Charger

<u>1.1.6</u> <u>Back of the Remote Display/Charger.</u> **See Figure 1-7** for the location of the major components on the back of the 3855 Remote Display/Charger.

- a. **Audible Speaker.** Provides the audible sounds for alarms and alerts.
- b. **2.4 GHz Antenna / Connector.** Used to connect 2.4 GHz Antenna for wireless communication with the 3850 MR IV pump.
- c. **Power Input.** Provides for connection to hospital grade AC Mains power.
- d. **Fuse Holder.** Provides replaceable 1A 3AG, 250V fuse for the AC Mains power.
- e. **Ground Terminal.** Used for grounding lug during electrical testing.
- f. **Battery Charger Compartment**. Located at the top rear of the Remote, allows charging of an 1133 Pump Battery when battery is installed and Remote is connected to AC Mains outlet as indicated by AC Power/Battery Charge LED (located below the power On/Off keys). Remote/Charger is AC powered only and will not run on battery power.

CAUTION: Battery pack is slightly magnetic. Use caution when transferring battery pack from pump to Remote/Charger near strong magnetic fields.



Figure 1-7. Back of 3855 Remote Display/Charger

1.2 Controls. The MRidiumTM 3850 MRI Infusion System is controlled with the use of "soft-touch" control keys. These control keys are located on the front panel of both the Pump and Remote Display. These control keys provide for turning the Pump and Remote Display ON or OFF, starting/stopping Channel's A and B infusion sequences, accessing and navigating in operational menus, review of data and setups, and for silencing any active alarms that may occur.

<u>1.2.1</u> Front Panel Control Keys. See Figure 1-8 for location of the control keys.



Figure 1-8. Front Panel Control Keys

START/STOP CHANNEL



START/STOP CHANNEL A. Pressing this control key starts, or stops, the Channel A infusion sequence.



CANCEL. Pressing the control key returns function to the previous menu or display with no action.



Up Arrow. Pressing this control key will increase the value of the currently selected option. It is also used for on-screen menu navigation.



Down Arrow. Pressing this control key will decrease the value of the currently selected option. It is also used for on-screen menu navigation.



MENU. Pressing this control key activates the Main Menu Display.



ENTER. Pressing this control key will activate, or select, a menu choice, proceed from one prompt to the next and accept/store changes.

START/STOP CHANNEL



START/STOP CHANNEL B. If the Channel B option (SideCar TM Pump mechanism) is installed, pressing this control key starts, or stops, the Channel B infusion sequence.



ALARM SILENCE. Pressing this control key temporarily silences audible alarms for two (2) minutes and clears alarms that have been resolved.



(1) Pressing this control key turns the unit on. When the unit is on this control key is inactive.



O Pressing and holding this control key for two (2) seconds turns the unit off.



Soft Keys. The six (6) Soft Keys, located to the left of the pump's display screen, are used to perform variable functions depending on the current state of the device. When active there is a small arrow present to the right of the key on the pump's display.

1.3 Display. The display screen provides the operator with the information and prompts necessary to operate this pump. See Figure 1-9 for a description of the Primary display. In general, the display screen functions as follows:

- **Highlighting.** As the different settings are "scrolled" through (using the Up or Down arrow control key), the selected item will become highlighted to distinguish it from the unselected items.
- Visual indicating of Soft key status. Active Soft keys have a "arrow" beside them pointing toward the menu item or setting that the Soft key controls. When Soft keys are inactive, the "arrow" is not displayed.
- **Split Screen.** The pump displays different types of information in the same area of the screen no matter what particular display is active.



Figure 1-9. Setup Display Screens

<u>1.3.1</u> <u>Informational Display.</u> The Informational Display is located at the top of the display screen. This portion of the display provides the operator with operational information such as the battery status, AC power status and the current time.



1.4

Provides a visual indication of the charge level of the Pump main battery. When full, the battery is charged fully. The level drops in increments of 25% on both the Pump and Remote as the battery discharges. If the symbol has an X through it no battery is installed.

Provides a visual indication of connection with Remote Display and receive level as indicated by vertical bars. Only displayed when internal radio has linked with Remote Display. Current channel is displayed to the right of the bars (1-6). **No bars indicates no communication**.



Provides a visual indication of the spare battery charge level in the Remote Display unit Informational Display. When full, the battery is charged fully. Level shows increments of 25% as the battery is charged, if symbol has an X through it no battery is installed.



Provides a visual indication that AC Main Power is connected on the Pump Informational Display.

06:11:55

Provides the current time in a 24 hour time format.

<u>1.3.2</u> <u>Infusion Parameter Setup Display.</u> Below the informational display, the Infusion Display provides windows into which the Volume to be Infused (VTBI) and the Infusion Rate are entered.

<u>1.3.3</u> <u>VI Display.</u> The VI Display provides the operator with the Volume Infused for the connected patient.

<u>1.3.4</u> <u>Secondary.</u> Pressing this soft key brings up the Secondary Infusion Setup Display to provide the operator with the ability to configure a secondary infusion.

1.3.5 <u>Bolus.</u> Pressing this soft key brings up the Bolus Setup Display to provide the operator with the ability to configure a Bolus infusion.



Figure 1-10. Special Features Setup Menu

<u>1.3.6</u> <u>Special Features Setup Menu.</u> Pressing the MENU button brings up the Special Features Setup Menu (See Figure 1-10) to provide the operator with the ability to set the Dose Rate Calculation, adjust the Alarm Volume, set the KVO Rate, adjust the Occlusion Limit, and Lock Keys and Next Menu if additional options are installed. The Lock Key will not be displayed unless it has been enabled in the Service Mode. (Refer to 1125 Service Manual for additional details on the Service Mode.)

1.4 User Interface. The user communicates with the unit by pressing the appropriate control and/or soft key and then using the up or down arrows and enter control keys to set and verify the setting being made.

1.5 System Self Test. The unit performs a series of power on Self-Tests on start-up to confirm readiness for use. Failure during any of these Self-Tests will result in a fault message and inability to use the pump. Verify successful completion of self-test prior to each use.

1.6 Maintenance and Operator Verification. (Using IVP1056 IV Set). Verify Pump and Channel B SideCar operation on a routine basis. Maintenance and verification must be performed at least once/year.

Test Flow Accuracy:	Flow pump output into a graduated cylinder with pump set at 200 mL/ hr. Run pump for 12 minutes and verify volume is 40 mL +/- 5%.
Test Inlet Occlusion:	With pump running 200 mL/hr close slide clamp on inlet side of pump and verify Inlet Occlusion Alarm.
Test Patient Occlusion:	With pump running 200 mL/hr close roller clamp on patient side of pump and verify Patient Occlusion Alarm.
Test Bubble Detector:	Insert a 100 uL bubble above the Bubble detector. Restart pump (ensure bubble flows down into detector), verify Bubble Detection Alarm.

Clean pump and inspect for any physical damage.

1.7 Cleaning Instructions. Clean Pump and Channel B SideCar and inspect for any physical damage after each use. Perform the following to clean the pump:

- a. Unplug power cord from AC outlet before cleaning.
- b. Remove Battery Pack from the rear of the Pump.
- c. Do not spray fluid directly into any connector.
- d. Use a soft cloth dampened with warm water and a mild, nonabrasive cleaning solution.
- e. A soft-bristled brush may be used to clean narrow areas.
- f. Use light pressure when cleaning pressure transducer and air-in-line detector areas of the pumping channels.
- g. Acceptable cleaning solutions include (*use per manufacturers' instructions*):
 - Soap and Warm water
 - Cidex[®] or other glutaraldehyde based surface disinfectants
 - Quaternary ammonium compounds
 - 10% Bleach Solution (1 part bleach to 9 parts water)

CAUTION: Do not use solvents or aromatic-solvent based cleaning agents. Damage to plastic parts of the Pump could occur. These include solutions containing aromatic solvents (naphtha, paint thinner, etc.), chlorinated solvents (Trichloroethane, Methy Ethyl Ketone (MEK), Toluene, etc.), alcohol, or phosphoric acid.

DO NOT use hard or pointed objects or pressurized sprays to clean any part of the Pump or Channel B SideCar.

DO NOT steam autoclave, EtO sterilize, or immerse the Pump or Channel B SideCar.

SECTION 2 INSTALLATION

2.0 Installation.

2.1 Introduction. This pump has been designed for the control and fluid management of patients undergoing magnetic resonance imaging.

2.2 Unpacking the Pump. Remove the pump and all accessories from their shipping carton and examine for visible damage that may have occurred during shipping. Check the materials carefully against the packing list and purchase request. Save all the packing materials, invoice and bill of lading as these may be required to process a claim against the carrier if there is damage from shipment. Contact Iradimed Corporation Customer Service for prompt assistance in resolving shipping problems.

The following is a list of items shipped with the pump:

- MRidiumTM 3850 MRI IV Pump.
- MRidiumTM 1124 Operations Manual.
- MRidiumTM 1125 Service Manual CD.
- Battery Pack, for MRidiumTM 3850 MRI IV Pump.
- Battery Charger/AC Adapter and Interconnect Cable.
- Hospital Grade Power Cord.
- One (1) Sample I.V. Set (Provided for initial Clinical Engineering testing).
- 2.4 GHz Antenna (with Remote Option)
- MRidiumTM 1127 Quick Reference Hang Tags

NOTE: Customer must order MRI infusion sets separately for use with this product.

2.3 Unpacking Remote Display/Charger. Remove the Remote/Display Charger and all accessories from their shipping carton and examine for visible damage that may have occurred during shipping. Check the materials carefully against the packing list and purchase request. Save all the packing materials, invoice and bill of lading as these may be required to process a claim against the carrier if there is damage from shipment. Contact Iradimed Corporation Customer Service for prompt assistance in resolving shipping problems.

The following is a list of items shipped with Remote Display/Charger:

- MRidiumTM 3855 Remote Display/Charger.
- Spare Battery Pack (Optional).
- 1129 Installation Guide.
- Hospital Grade Power Cord.
- 2.4 GHz Antenna.
- LB2025 Channel ID Marker Sheet

2.4 Unpacking 3851 Channel B SideCar. Remove the Channel B SideCar from it's shipping carton and examine for visible damage that may have occurred during shipping. Check the materials carefully against the packing list and purchase request. Save all the packing materials, invoice and bill of lading as these may be required to process a claim against the carrier if there is damage from shipment. Contact Iradimed Corporation Customer Service for prompt assistance in resolving shipping problems.

The following is a list of items shipped with Channel B SideCar:

- MRidiumTM 3851 Channel B SideCar.
- One (1) Sample I.V. Set (Provided for initial Clinical Engineering testing).

• 1130 Installation Guide

2.5 Preparing the Pump for Use. Once the pump has been unpacked, perform the following to prepare the pump for use.

<u>2.5.1</u> <u>Installing Battery.</u> Slide the removable battery pack into the rectangular open slot on the rear of the MRidiumTM 3850. The battery pack will lock into place.

a. **Battery Initial Charging.** The battery should be allowed to charge for nine (9) hours prior to placing the pump into use. The battery charges whenever the pump is connected to a hospital grade power outlet through the AC Power Adapter.

CAUTION: Use only non-magnetic I.V. pole designed for proper safe support of the MRidiumTM 3850 I.V. pump.

2.6 Mounting on I.V. Pole. The Pump should be mounted on an MRI-compatible (nonmagnetic) I.V. pole and secured with its ergonomically designed clamp knob. The clamping mechanism accommodates pole diameters 1 to 1.5 inches (25 to 38 mm).

WARNING: When using multiple Pumps (limit 2), distribute the Pumps to ensure stability.

2.7 Operational Checkout of the Pump. Power up pump as follows and confirm no failures are displayed during the Power Up Test.

- a. Turn on pump.
- b. Observe while pump performs power up test.
- c. Verify that the pump sounds an audible beep.
- d. Observe that the pump displays the initial start-up screen.

2.8 Pump Storage. Plug the Pump into an AC outlet during storage to ensure a fully charged battery when needed. The AC indicator light will be green or amber (if charging) whenever Pump is plugged in to the AC Power. Close the door latch(es) whenever the Pump or Channel B SideCar are not in use.

2.9 Remote Display Installation. See Paragraph 1.1.6. Connect 2.4 GHz antenna and AC Power Cord to Remote Display/Charger. Verify MRidiumTM 3850 MRI IV Pump has Remote Radio Option installed. And perform the following:

NOTE: Information shown on the remote display is updated at least once per second when communicating with the pump.

- a. Turn both Pump and Remote Display/Charger on.
- b. Press the Menu key on the Remote Display/Charger
- c. See "NEXT MENU key" on page 4-17. Press the NEXT MENU soft key to bring up additional menu options, then press the Set Comm Channel soft key.
- d. From the Radio Channel menu, select the desired Channel 1 through 6 by pressing the soft key next to that channel and verify that Channel is now highlighted.
- e. Press the Menu key on the MRidiumTM 3850 MRI IV Pump.
- f. **See "NEXT MENU key" on page 4-17.** Press the NEXT MENU soft key to bring up additional menu options, then press the Set Comm Channel soft key.
- g. From the Radio Channel menu, select the same Channel that was selected for the Remote by pressing the soft key next to that channel and verify that Channel is now highlighted.

- h. Apply an ID marker to both the Remote and Pump indicating the Channel selected. Do not apply an ID Marker to the Channel B SideCar module as it can be removed to use on another pump.
- i. Verify communication between Remote and Pump by pressing the Menu key on one and the Cancel key on the other, and observing both display match and change simultaneously.

WARNING: Never Set two Pumps or two Remotes on the same Comm Channel. Always confirm the Remote Display is communicating with the selected Pump before use.

<u>2.9.1</u> <u>Charging Battery with the Remote.</u> If a spare battery is to be charged (See Figure 1-7), insert battery into the Remote Display/Charger's battery bay.

Verify spare battery Icon is visible on the Remote Display/Charger's informational display area. The AC/Power LED on the bottom, left-hand side will be amber colored until the battery is fully charged. The LED will turn green when battery is fully charged and the spare battery Icon in the Remote Display/Charger's informational display area will be full with 4 bars.

NOTE: If communication between Remote and Pump is lost the Remote will sound a low pitched alarm for approximately 10 seconds. If communication is not returned after 10 seconds the Remote will sound a ramping high pitch alarm tone and display "No Communications."

2.10 Language Options. The MRidium System is capable of displaying information in languages other than English. This feature is service selectable, and should only be changed by qualified service personnel, as described in the 1125 Service Manual.

2.11 Product Structure.The MRidium 3850 MRI Infusion system is mainly consisted of the following:



3850R (Pump Main Body)



3851 (with 3850R) SideCar Pump



3855 Remote Display/Charger



1120 Power Supply



1119 IV Pole Stand
SECTION 3

I.V. SET PREPARATION FOR USE

3.0 I.V. Set Preparation for Use.



Figure 3-1. Free Flow Preventer

(Free Flow Preventer is shown in the no flow/closed position).

a. **Operation of the Free Flow Preventer.** (See Figure 3-1) The Free Flow Preventer is used to stop the flow of the fluid through the infusion set. The Free Flow Preventer will open automatically to allow fluid flow when the pump door is shut and will also close automatically to prevent fluid flow when the pump door is open.

WARNING: To prevent free flow, verify Free Flow Preventer is in the closed position (slide clamp pulled out). When properly loaded in the pump, the Free Flow Preventer in the infusion set automatically prevents free flow when the pump door is opened. It can also be manually operated for user set-up convenience. The "CHECK DOOR" alarm indicates Free Flow Preventer is in the 'open' flowing position with the pump door not fully closed. Immediately check for proper IV set installation, Free Flow Preventer position, and pump door closure.

b. **Operation of the Roller and Slide Clamps.** Either the Roller Clamp (See Figure 3-2a), or the Slide Clamp (See Figure 3-2b), is the manual way used to stop the flow of the fluid through the infusion set. Close the Roller Clamp by moving the roller to pinch the tubing closed, or close the Slide Clamp by moving the clamp to pinch the tubing closed.



Figure 3-2. Roller and Slide Clamps

- **3.1 Precautions.** The following precautions apply to the use of these I.V. Sets:
 - Always use aseptic technique when handling any I.V. Set.
 - The fluid path is sterile and nonpyrogenic, discard if packaging is not intact or if protector caps are missing or loose.

- Replace Set according to accepted hospital policies or every six (6) hours maximum.
- For I.V. push medication, occlude tubing above injection port.
- Contraindicated for most blunt cannula systems.
- For unattended medication delivery; only luer-locking Secondary sets and syringes should be attached to the valve.
- Flush injection ports in accordance with hospital policy after or between: small volume injections; lipids; blood or blood product infusions; blood withdrawals; incompatible medications.
- FREE FLOW-PREVENTER is OPEN when half-moon shaped clamp is pressed INWARD. To CLOSE, PULL the slide clamp OUTWARD.
- Dispose of used set in accordance with applicable regulations and hospital policy.
- Follow appropriate pump directions for use.

3.2 I.V. Set Priming. There are four I.V. Sets available for use with this pump. Priming of Administration Set 1056 is discussed in subparagraph 3.2.1; priming of ByPass Set 1055 is discussed in subparagraph 3.2.2; priming of Syringe Adapter Set 1057 is discussed in subparagraph 3.2.3. and priming of Extension Set 1058 is discussed in subparagraph 3.2.4. Always use aseptic technique.

<u>3.2.1</u> <u>I.V. Administration Set 1056 Priming.</u> Refer to instructions accompanying the set. Perform the following to prime I.V. Administration Set 1056:

- a. Remove from package, close roller clamp and remove blue tubing protector from silicone rubber pumping segment.
- b. Insert administration set spike into prepared fluid container following accepted hospital procedure and hang fluid container approximately 15 inches above MRidiumTM 3850 Pump.
- c. Squeeze drip chamber to fill approximately 1/2 full.
 - Open vent cap if glass fluid container.
- d. To prime tubing and clear air from injection sites, slowly open roller clamp, allowing fluid to fill tubing.
- e. Invert and tap injection sites to clear any air bubbles.
- f. Close roller clamp and Free Flow Preventer by sliding pinch clamp outward.

WARNING: Inlet Occlusion indication may not be reliable when used with bypassed I.V. sets that incorporate a check valve.

<u>3.2.2</u> <u>I.V. ByPass Set 1055 Priming.</u> Refer to instructions accompanying the set. Perform the following to prime I.V. ByPass Set 1055:

- a. **Ensure hospital pump set tubing is clamped closed**. Remove set from hospital pump.
- b. Remove ByPass set from package, close roller clamp and remove blue tubing protector from silicone rubber pumping segment.
- c. Attach clear covered Luer Lock connector to upper injection site on hospital set.
- d. Open roller clamp on ByPass set, allow fluid to fill tubing.
- e. Tap injection sites to clear any air bubbles.
- f. Close roller clamp, close Free Flow Preventer pinch clamp by sliding it outward.
- g. Attach blue capped Luer lock connector to lower injection site on hospital set.

<u>3.2.3</u> <u>I.V. Syringe Adapter Set 1057 Priming.</u> Refer to instructions accompanying the set. Perform the following to prime I.V. Administration Set 1057:

NOTE: Do not puncture or add medication through air inlet (See Figure 3-3f).

- a. Remove from package, close slide clamp and remove blue tubing protector from silicone rubber pumping segment.
- b. Remove protective Clear Cap. Attach syringe with fluid to vented Syringe Adapter Fitting at top of I.V. set. Follow accepted hospital procedure. Ensure Tube Slide Clamp and FLOW-PREVENTER are open.

NOTE: When attaching syringe, ensure air vent tube is placed inside the syringe fluid path opening before luer-locking the syringe.

- c. Manually push syringe to prime I.V. Set removing all air bubbles.
- d. Open air vent cap on Syringe Adapter Fitting (See Figure 3-3e.)
- e. Close slide clamp, and close FLOW-PREVENTER by sliding black pinch clamp outward.
- f. Grasp FLOW-PREVENTER slide clamp and load pump segment into Pump.
- g. Close Pump Door. Ensure syringe with fluid is mounted vertically after set is loaded and Door is closed.
- h. Attach distal luer-outlet (White cap) to the port nearest the patient's vascular access device.
- i. Open set slideclamp and begin infusion.

CAUTION: Do not strike or jar the syringe adapter it has been loaded onto the 1057 Syringe Adapter Set and mounted onto the pump. I.V. set or syringe damage could occur.

NOTE: Small diameter tubing (0.040 inch) is used in the 1057 Set. Dependent on the desired Flow Rate and fluid viscosity, the Pump's Occlusion Pressure Limit may need adjustment to avoid false alarms. Pressures may exceed the maximum allowable limit for Flow Rates above 600 mL/hr.

Do NOT depress the syringe plunger during infusion. Fluid leakage out the air inlet can occur. Since this set is vented, fluid will flow without movement of the syringe plunger. If the patient's access device requires needle access, add a 17 Gauge (or larger) needle to the 1057 set's luer-outlet before attaching to the access device.

<u>3.2.4</u> <u>I.V. Extension Set 1058 Priming.</u> Refer to instructions accompanying the set. Perform the following to prime I.V. Extension Set 1058:

- a. **Ensure hospital pump set tubing is clamped closed.** Remove pre-existing infusion set from its non-MRI hospital pump. Ensure flow is stopped/clamped-closed before removal.
- b. Remove 1058 Extension set from package, close both the roller clamp and FLOW PREVENTER and remove blue tubing protector from silicone rubber pumping segment.
- c. Attach inlet luer-fitting (clear cap) to the pre-existing infusion set at its distal end.
- d. Open roller set clamp to prime the set. Ensure FLOW PREVENTER is open.
- e. To remove air from the injection port, invert and tap while fluid is passing through the port.

- f. Close FLOW PREVENTER pinch clamp by pulling the black slide clamp outward.
- g. Grasp FLOW PREVENTER slide clamp and load pump segment into MRidium pump.
- h. Attach distal luer-outlet (blue cap) to pre-existing infusion set's injection port nearest the patient's vascular access device.
- i. Open set roller clamp and begin infusion.

3.3 I.V. Set Insertion and Removal. For insertion instructions see subparagraph 3.3.1 and for removal instructions see subparagraph 3.3.2.

WARNING: DO NOT STRETCH the silicone rubber segment of the I.V. Set. When properly loaded, the silicone rubber segment should be taut (without slack) and centered atop the pump's peristaltic "fingers", as shown in **Figure 3-3**, page 3-5. Overstretching, or mispositioning of this tubing can result in inaccurate flow, and/or free-flow.

WARNING: Confirm that the I.V. Set is fully primed prior to inserting the set into the pump.

<u>3.3.1</u> <u>Administration Set Insertion.</u> Perform the following to insert the Administration set into the pump: (See Figure 3-3)

- a. Press the ON Control Key (I).
- b. Open MRidiumTM 3850 Pump door.
- c. Install administration set pumping segment by properly positioning the upper tubing retainer (see Figure 3-3a) and ensure Free Flow Preventer is closed (see Figure 3-3b).

DO NOT STRETCH TUBING

- d. To insert the Free Flow Preventer, tilt the Free Flow Preventer, top in first (see Figure 3-3c), and rotate bottom downward to lock into place using Free Flow Preventer's slide clamp. Assure tubing above Flow Preventer is located directly down the middle of white PUMP "fingers" (see Figure 3-3g).
- e. Run tubing through the Air-in-Line detector (see Figure 3-3d) at the bottom of the pump.
- f. Close MRidiumTM 3850 Pump door.
- g. Open roller clamp, or slide clamp, depending on IV set type.
- h. Verify no flow is flowing through the drip chamber (1055, 1056, or 1058 type IV Sets), or if using 1057 syringe set, be sure to open vent cap prior to starting pump (see Figure 3-3e-f).

<u>3.3.2</u> <u>Administration Set Removal.</u> Perform the following to remove the Administration set from the pump:

- a. Press the START/STOP Channel control key on the MRidiumTM 3850 Pump
- b. Close roller clamp, or slide clamp, depending on IV set type.
- c. Open the MRidiumTM 3850 Pump door, which automatically closes the Free Flow Preventer
- d. Remove the tubing by grasping the Free Flow Preventer by the Slide Clamp and push down while tipping the top outward.

The tubing will allow gravity flow when the Free Flow Preventer pinch clamp and the roller clamps are in the open position. This provides the ability to infuse fluids/medications via gravity in an emergency situation.

WARNING: Check Free Flow Preventer (black side clamp) is fully pulled outward to the shut off position.

CAUTION: Check Free Flow Preventer (black side clamp) is fully pulled outward (shut off) any time the chamber door is not fully latched in the closed position. Partial closure of the door may open the Free Flow Preventer, always close door and fully latch.



Figure 3-3. IV Set Installation

SECTION 4 PUMP PREPARATION FOR USE

4.0 **Pump Preparation for Use.**

4.1 Quick Setup. Perform the following to setup the pump:

Press the (1) control key to power on the MRidiumTM 3850 Infusion Pump.

- a. If displayed, select "Same Patient" to use the last infusion settings or "New Patient" for new dose information. This menu will not be displayed if pump is off for more than one (1) hour.
- b. Adjust Rate using the Up and Down arrow keys, then press the **EVER** control key.
- c. Adjust VTBI using the Up and Down arrow keys, then press the **ENTER** control key.
- d. Open I.V. Set Compartment door and install primed and de-bubbled MRidiumTM 1000 Series I.V. Set. Take care that the Black Free Flow Preventer tab is pulled outward.
- e. Shut the I.V. Set Compartment door and press the Start/Stop control key for the desired pump channel when the "Ready to Start" message appears.
- f. Monitor pump is correctly delivering I.V. therapy by viewing the green blinking light above the selected channel. Check fluid drip rate in the drip chamber.

WARNING: This pump will stop fluid flow during Alarm Conditions. Periodic patient monitoring must be performed to ensure that the infusion is operating normally.

WARNING: It is the responsibility of hospital personnel to ensure that drugs used in this system are compatible as well as ensuring the performance of each pump as part of the overall infusion. Potential hazards include, but are not limited to, drug interactions, inaccurate delivery rates, inaccurate pressure alarms and nuisance alarms.

WARNING: The use of positive displacement infusion devices connected together with gravity flow infusion systems into a common I.V. site may impede the flow of common "gravity only" systems and affect their performance. It is the responsibility of hospital personnel to ensure that the performance of the common I.V. site is acceptable under the circumstances in which it is being used.

<u>4.1.1</u> <u>Administration Sets.</u> Use only Iradimed Corporation approved MRidiumTM 1000 series infusion sets. The use of any other set will cause improper pump operation and will result in inaccurate fluid delivery and risk to the patient.

Before beginning any infusion cycle, verify that the infusion lines are free from kinks and are loaded correctly on the pump.

Do not reuse any component that is labeled for single use. Discard these items immediately after use.

<u>4.1.2</u> <u>Artifacts.</u> It is normal for this pump to produce non-hazardous currents when being used for the infusion of electrolytes. These currents will vary proportional to the pump's infusion rate. Therefore, when an ECG monitoring system is not functioning under optimal conditions, these currents may appear on the ECG display screen as artifacts and simulate actual ECG readings. To determine if the ECG abnormalities are caused by patient condition or the ECG equipment receiving artifact from the pump, place the infusion pump on HOLD and observe the ECG signal. If the ECG signal becomes normal, the ECG monitor requires attention as proper setup of the ECG equipment should remove the pump caused artifact. Reference the appropriate ECG monitoring system documentation for instructions on setup and maintenance of the ECG device.

NOTE: Operating parameters (Power, VTBI, etc.) are retained in memory for one (1) hour unless all power is lost (no AC and a depleted battery). A "Settings Lost" message at power up indicates existing operating parameters have been erased, and the system has reverted back to initial default parameter settings.

The operator may choose "New Patient" or "Same Patient." Same patient will pre-set primary pump setup display with the prior Rate and VTBI. New patient will clear the Rate, VTBI and VI.

If the Primary Pump Setup screen does not appear as shown in Figure 4-1, it may indicate improper functioning of the display. Although the Pump may appear to function properly, return it to qualified service personnel for examination and/or repair.

CAUTION: Each time the Pump is turned on verify and/or set the pressure alarm limit. If the pressure alarm limit is not verified, the Pump may not be operating with the desired occlusion detection parameter(s).

4.2 Primary Configuration. Pressing the <u>III</u> control key powers up the pump. Upon power up the pump first runs a Power On Self-Test then displays the message "Powering On" and enters the Primary Pump Setup screen (See Figure 4-1) where the Infusion Rate and Volume to be Infused (VTBI) is set.

The pump memory will retain the last used settings for up to one (1) hour following power down of the pump). Operator has the option of selecting either "New Patient" or "Same Patient."

If any part of the Power On Self-Test fails, the pump displays an error message and will not operate.

If optional Channel B SideCar is installed on the left side of the display, the user may select and program Channel B in the same manner as Channel A by selecting the Channel B softkey.



Figure 4-1. Primary Pump Setup Screen

<u>4.2.1</u> <u>Programming a Basic Infusion.</u> Perform the following to enter the primary Rate and VTBI for infusion:

- a. Insert primed MRidiumTM 3850 I.V. Set into pump and close door.
- b. Observe that the RATE value is highlighted (if not highlighted, select using the soft key to the left of Rate).
- c. Press the up and down arrow control keys to enter the desired flow rate.
- d. Press ENTER or VTBI soft key to move to the next field.
- e. Observe that the VTBI value is highlighted.
- f. Use the up and down arrow control keys to enter the desired volume to be infused.
- g. Observe that the Ready to Start Prompt is displayed. If Ready to Start is not displayed, check the I.V. Set loading and for proper door closure/latching.

NOTE: The VTBI is displayed on the main display screen of the MRidiumTM 3850 pump and will count backwards to 0 mL. At that time the infusion rate will change to the predetermined KVO rate, and the VTBI will display KVO rate.

- h. Press the START/STOP (A or B) Channel control key for the proper Channel to begin the infusion.
- i. Observe that the rate and VTBI are displayed and the green infusing indicator is flashing.

j. Verify infusion in running from primary container by observing the drip chamber, or air bubbles within the loaded syringe (1057 set only).

NOTE: The Pump needs to be ON prior to loading the I.V. Set. If tubing is installed prior to turning Pump on, "Load Set" or "Unload Set" messages may appear. Remove and replace I.V. Set to clear. "Ready to Start" prompt will appear.

<u>4.2.2</u> <u>Editing the Infusion Program.</u> The clinician has the ability to change or edit (titrate) the infusion program without having to pause the infusion. Perform the following to change or edit the infusion program:

- a. Press the RATE or VTBI soft key and set the new parameter(s) using the up and down arrow control keys.
- b. Press ENTER to accept the new value.

NOTE: While the pump is in use, all infusion parameters are stored in memory unless the pump has been powered down for more than one (1) hour.

<u>4.2.3</u> <u>Canceling/Pausing an Infusion.</u> The clinician has the ability to pause an infusion by pressing the START/STOP Channel control key (for the channel to be cancelled/paused) once. When the START/STOP Channel control key has been pressed, the pump will display CANCEL INFUSION next to a soft key and the indicator light will go out. Every 10 seconds, the pump will give a periodic audible tone to remind the operator that the pump is still on hold.

- a. To cancel the infusion, press the soft key next to the CANCEL INFUSION message.
- b. To restart the infusion, press the START/STOP Channel control key for the channel to be restarted.

<u>4.2.4</u> <u>KVO (Keep Vein Open) - Infusion Complete.</u> During an infusion, the VTBI value will decrease until the VTBI reaches 0 mL. When the VTBI is 0 mL, the infusion automatically switches to the configured KVO rate, or remains at the programmed primary flow rate, whichever is less. KVO is displayed next to the VTBI and the RATE will display either the preselected KVO rate or the current infusion rate (again, whichever is less). A short tone will announce the switch to KVO. The KVO rate is configurable from the Special Features Menu (which is accessed by pressing the MENU control key) and has a rate range from 1 mL/hr to 5 mL/hr.

- a. **Resuming an Infusion after KVO.** Perform the following to resume an infusion after KVO has initiated:
 - (1) Press the START/STOP Channel control key.
 - (2) Replace infusion bag/bottle.
 - (3) Press the VTBI soft key.
 - (4) Observe that the VTBI value is highlighted.
 - (5) Use the up and down arrow control keys to enter the new volume.
 - (6) Observe that the Ready to Start prompt is displayed.
 - (7) Press the START/STOP Channel control key to resume infusion at the prior rate.

<u>4.2.5</u> <u>Volume Infused.</u> The MRidiumTM 3850 pump tracks the Volume Infused (VI) when the device is used to deliver fluid to the patient. The VI on the device is cumulative with the system calculating the total VI including both the primary and secondary infusions. The operator has the ability to clear the VI for a specific channel or for all attached channels from the setup screen only. The VI is displayed on the run and setup screens.

- a. **Clearing Total Volume Infused.** Perform the following to clear the total Volume Infused from the setup screen:
 - (1) Press the VI soft key.
 - (2) Observe that displays the message "Press VI again to clear display."
 - (3) Press the VI soft key again to clear the total volume infused.

<u>4.2.6</u> <u>Pump Shut Down.</u> When the pump is powered down it will retain the infusion parameters (including volume infused) and operational setups for a period of one (1) hour. Perform the following to shut down the pump:

- a. If an infusion is in process press the START/STOP Channel control key to place the infusion on hold then press the **Cancel Infusion** soft key.
- b. Press and hold the i control key for two (2) seconds, an audible tone will sound then the pump will power down.

<u>4.2.7</u> <u>Restoring an Infusion following Pump Power Down.</u> Perform the following to restore an infusion after shutting down the pump:

- a. Press the 💷 control key.
- b. Observe that "Resume Therapy" appears after the self check, if it has been less than one (1) hour since shut down. The display will prompt for the selection of either "New Patient" or "Same Patient." Press the soft key to select the appropriate choice. (If power has been off for more than one (1) hour, the primary setup screen will display "New Patient" only).
- c. Press the Rate soft key.
- d. Observe that the rate value is highlighted.
- e. Adjust the Rate using the up and down arrow control keys if necessary.
- f. Press Enter to accept value.
- g. Observe that the VTBI value is highlighted.
- h. Use the up and down arrow control keys to change the VTBI value if necessary.
- i. Press the VI soft key twice to clear the volume infused if necessary.

4.3 Dual Channel Infusion. The MRidiumTM 3850 pump may be configured to perform simultaneous infusions on both Channel A and B (if equipped with the optional SideCar).

<u>4.3.1</u> <u>Dual Channel Infusion Setup.</u> To perform a dual channel infusion, configure both Channel A and Channel B as described in Paragraph 4.2. Push the Start/Stop buttons of both channels to begin the infusion. Once both infusions are started, the display screen "splits" to display information for both infusion channels (Figure 4-2).



Dual Channel Infusion Split Screen

Figure 4-2. Dual Infusion Display

4.4 Secondary Infusion Configuration. The MRidiumTM 3850 pump supports secondary infusions in the form of automatic sequential piggybacking. Medications must be compatible to be infused in this manner, as they will mix in the tubing below the injection site. When the device is programmed and delivering in the secondary mode, only fluid from the secondary container is being delivered to the patient as the primary infusion is temporarily stopped. Delivery from the primary container resumes when the fluid level in the secondary line equilibrates with, or is even with, the fluid level in the primary container.

There are key elements in setting up and delivering a secondary infusion.

- Head height differential. To prevent sympathetic flow (flow from the primary container), the secondary container must be at least 9.5 inches (24 cm) higher than the primary container.
- The clamp on the secondary set must be opened. If the clamp is not opened, the fluid will be delivered from the primary container.

WARNING: The secondary VTBI settings require consideration of such variables as factory overfill, medication additions, etc. Underestimating the volume will cause the remaining Secondary solution to be infused at the Primary rate; overestimating will result in the Primary solution being infused at the secondary rate. Multiple doses from a single container are not possible.

WARNING: The Secondary set must be primed prior to beginning the Secondary infusion.

<u>4.4.1</u> Priming a Secondary Administration Set. Any secondary set can be used with the MRidiumTM 1000 Series I.V. Set. The secondary set must be primed prior to beginning the secondary infusion. The secondary set may be primed in one of two ways: 1.) Forward Priming: Gravity prime using the secondary fluid, or 2.) Back Priming: Priming the secondary line using the fluid from the primary container (the back priming technique minimizes the amount of wasted medication that might occur using the Forward Priming technique).

NOTE: The Back Priming technique may be used for subsequent doses of medication.

If there is any question on drug incompatibilities, consult with the hospital pharmacist before performing the Back Prime technique.

- a. **Back Priming Technique.** Perform the following to back prime a secondary administration set (using aseptic technique):
 - (1) Spike secondary medication bag and clamp secondary tubing.
 - (2) Attach the secondary set to the upper injection port on the primary administration set.
 - (3) Lower the secondary fluid container below the upper injection port on the primary administration set.
 - (4) Open the clamp on the secondary set to allow fluid from the primary to "back prime" into the secondary line
 - (5) When the secondary set is completely primed, close the clamp and suspend the secondary container on the I.V. pole higher than the primary fluid container.
- <u>4.4.2</u> <u>Secondary Infusion Setup</u> Perform the following to setup a secondary infusion :
 - a. Lower the primary fluid container onto a non-magnetic hanger (may be enclosed with the secondary administration set). Verify hanger is non-magnetic prior to use. This will allow the pump to pull from the secondary fluid container.
 - b. Attach the primed secondary set to the upper injection port of the primary administration set or back prime as above.
 - c. Open the clamps on both the primary and secondary administration sets.
 - d. When the fluid level in the secondary line equilibrates with the fluid level in the primary container, flow from the primary fluid container resumes.



Figure 4-3. Secondary Infusion Set-up and Programming Screen

NOTE: The Channel B option will only appear when an optional second pump channel is installed.

<u>4.4.3</u> <u>Programming a Secondary Infusion.</u> Perform the following to program a secondary infusion:

- a. Press the SECONDARY soft key.
- b. Observe that the programming screen splits to display Secondary Rate and VTBI with the Rate value highlighted (See Figure 4-3).
- c. Enter the desired flow rate using the up and down arrow control keys.
- d. Press Enter or VTBI soft key.
- e. Observe that the VTBI value is highlighted.
- f. Enter the desired volume to be infused using the up and down arrow control keys.
- g. Observe that the Ready to Start prompt is displayed.
- h. Press the START/STOP Channel control key.
- i. Observe that the secondary Rate and VTBI is displayed on the main display screen.
 - The green infusion running indicator light flashes.
 - Confirm flow from the secondary fluid container by observing the drip chamber.
 - When the VTBI = 0 mL, the pump gives an audible alarm and the primary parameters resume and are displayed.

WARNING: The clamp on the secondary set must be opened. If the clamp is not opened, the fluid will be delivered from the primary container.

The Secondary VTBI must be equal to the volume in the secondary container. Variables such as factory overfill and the amount of medication must be considered when programming the VTBI.

Underestimating the secondary VTBI will result in the remainder of the secondary solution to be delivered at the primary rate.

<u>4.4.4</u> <u>Viewing Primary Settings during a Secondary Infusion.</u> It is possible to view the primary settings while a secondary infusion is in process. Perform the following to view the primary settings during a secondary infusion:

- a. Press the Adjust Primary soft key.
- b. Observe that the primary parameters are now displayed on the central display screen with the secondary infusion continuing without interruption.
- c. Press the CANCEL soft key to return to the secondary run screen (or wait five (5) seconds for Timeout Return).

<u>4.4.5</u> <u>Changing Primary Settings during a Secondary Infusion.</u> It is possible to change the primary settings while a secondary infusion is in process. Perform the following to change the primary settings during a secondary infusion:

- a. Press the Adjust Primary soft key.
- b. Observe that the Primary parameters are displayed with the Rate value highlighted.

- c. Press the RATE or VTBI soft key and set the new parameter(s) using the up and down arrow control keys.
- d. Press Enter to accept the new values and return to Secondary display, or press the CANCEL soft key to return to the secondary run screen.

<u>4.4.6</u> <u>Stopping a Secondary Infusion and Returning to the Primary Infusion</u>. Perform the following to stop a secondary infusion and return to the primary infusion:

- a. Press the START/STOP Channel control key.
- b. Observe that **Cancel** (A) or (B) is displayed as a soft key option.
- c. Press Cancel (A) or (B) soft key.
 - The pump is on hold and the light is not on.
- d. The Primary infusion programming screen will display.
 - To stop flow from the secondary container, the clamp on the secondary set must be closed or the set disconnected from the upper injection port.
- e. Press START/STOP to restart at primary settings.

<u>4.4.7</u> <u>Dual Channel Operation.</u> When both channel A and B are running, the options for viewing and changing the primary settings when a secondary infusion is running move to the "Menu". To view or change the primary settings:

- a. Select MENU Key.
- b. Press Primary A or Primary B Soft Key
- c. Verify that Primary A Rate is highlighted. Use UP or DOWN arrow keys to change if necessary.
- d. Press VTBI Soft Key. Use UP or DOWN arrow keys to change if necessary. Press ENTER to save changes and return o run screen.
- e. Press CANCEL to return to split running screen without making changes.





4.5 Bolus Delivery. The Bolus feature is used to deliver a bolus. The operator enters this feature by selecting the Bolus option in the menu on the setup screen or the running screen.

<u>4.5.1</u> <u>Bolus Setup and Start.</u> The Bolus option may be programmed at the start of an infusion or added to an infusion in progress. Perform the following to program a Bolus Dose for infusion:

NOTE: The VTBI of the Bolus cannot exceed the VTBI of the Primary infusion.

- a. **Bolus A Soft Key:** If the Channel B SideCar Option is installed and Channel A channel is running pressing this soft key will allow Bolus A selection.
- b. **Bolus B Soft Key:** If the Channel B SideCar Option is installed and Channel B channel is running pressing this soft key will allow Bolus B selection.
- c. Press BOLUS soft key from either the Primary programming, running screen or menu.
 - If from the primary programming screen the screen will split for Bolus information (See Figure 4-4).
 - If from the primary running screen, a separate Bolus display will appear (See Figure 4-4).
 - When both channels are running, the Bolus option is accessed from the "Menu." The screen will display "Bolus" for the channel chosen.
- d. Observe that the Rate is highlighted.
- e. Use the up and down arrow control keys to enter Bolus rate (the rate defaults to 500).
- f. Press the VTBI soft key.
- g. Use up and down arrow control keys to enter Bolus VTBI value.
- h. Press Enter to accept (this starts the Bolus if programmed while the pump is running).
- i. If pump was not running, press the START/STOP Channel control key to begin Bolus infusion.
- j. Upon completion of the Bolus infusion, an audible alert sounds and the primary infusion rate resumes with the VTBI counting down.
- <u>4.5.2</u> <u>Stopping Bolus Rate Dose.</u> Perform the following to stop a Bolus Dose infusion:
 - a. Press START/STOP Channel control key
 - b. Press Cancel (A) or (B) soft key.
 - c. Observe that the **Push Start** (**A**) or (**B**) prompt is displayed.
 - d. Press the START/STOP Channel control key to resume the Primary infusion parameters.

<u>4.5.3</u> <u>Restoring Bolus Dose.</u> A Bolus Dose can be restored after it is complete. Perform the following to restore a Bolus Dose:

- a. Press BOLUS soft key from running screen or Menu.
- b. Verify dosing parameters and press Enter control key. The Bolus begins immediately.

NOTE: Confirm the primary mode parameters are correct before accessing the Bolus Dose option. Bolus VTBI must be less than the primary VTBI.





Figure 4-5. Attaching Channel B "SideCar"

4.6 Channel B. (3851 I.V. Pump SideCarTM) A second channel may be added to the MRidiumTM pump. Perform substep **4.6.1** to add the Channel B SideCar and substep **4.6.2** to remove the Channel B SideCar.

<u>4.6.1</u> <u>Attaching the Channel B SideCar.</u> Perform the following to attach the Channel B SideCar:

- a. Turn 3850 Pump OFF.
- b. Remove SideCar port protector cap on rear of 3850 Pump. (Do not discard protector cap).
- c. Position the Channel B SideCar (See Figure 4-5) next to the left side of MRidiumTM 3850 Pump
- d. Slide the Channel B SideCar connection port onto the pump, press together and tighten two (2) thumb screws on the rear. Tighten upper and lower screws a little at a time until secure. Do not overtighten the thumbscrews.
- e. Turn ON pump and verify start-up checkout (refer to section 4.2).

<u>4.6.2</u> <u>Detaching the Channel B SideCar.</u> Perform the following to detach the Channel B SideCar:

- a. Turn 3850 Pump OFF.
- b. Slide the Channel B SideCar backward to disconnect from main pump after loosening upper and lower thumb screws.
- c. Re-attach SideCar port protector cap on rear of 3850 pump.

4.7 Special Features Menu. There are additional features that are accessed by pressing the MENU control key. Pressing the MENU control key brings up the Special Feature Setup Menu to provide the operator with the ability to set the Dose Rate Calculation, adjust the Alarm Volume, set the KVO Rate and adjust the Occlusion Limit. Depending on optional accessories (i.e. Remote or Channel B) user may also lock keyboard, set communication channel for Remote Display, program Bolus or change/view primary infusion from this screen.



Figure 4-6. Special Feature Setup Menu



d earlier and above Figure 4-7. Dose Rate Calculation Menu <u>4.7.1</u> <u>Dose Rate Calculation.</u> The MRidiumTM 3850 infusion pump automatically calculates a Rate based on a Dose amount, Weight and Concentration of medication. The correct rate and dose is then automatically transferred to the main programming screen (for Dose Rate calculations and Drug Library included in software releases later than Release 362, see chart below and Default Values Table in Appendix A on page A-3).

Dose Rate Calculator w/Drug Library (Only available in software releases later than Software Release 362). Dose Rate Calculator can be used for Primary, Secondary or Bolus infusions.		
 A. Drug Library 1. Drug? (Default) 2. Propofol 3. Dobutamine 4. Adenosine 5. Dexmedetomidine 	 Dose Rate Calculator Default Values Table provided in Appendix A on page A-3. WARNING: Selection of any drug from the Dose Rate Calculator Library will replace all user set values to the default values described in the Default Values Table in Appendix A. Verify all selections prior to beginning the infusion. 	
 B. Dosing Units Original Dose Units: mg/kg/min mg/kg 2. Volume over Time Dos mcg/kg 0.00 - mg/kg 0.00 - 	y/hr mcg/kg/min mcg/kg/hr se Units: 3. Non Weight Based Dose Units: 9999 mg/hr mg/min 9999 mcg/hr mcg/min 9 min No Weight Entry	
 Messages are developed to indicate if calculated values exceed the Rate or VTBI specifications: Und (under-range) or Ovr (over-range) denote settings outside the established units: Rate: 1.0 to 99.9, 100 to 999 mL/hr VTBI: 0.1 to 999 mL NOTE: In Pediatric Mode, the Dexmedetomidine unit values are fixed at Dose = mcg/kg/hr and CONC = mcg. These unit values can only be modified in the Adult Mode. NOTE: Drug Library can be enabled/disabled from the Service Mode. This feature should only be set by qualified service personnel, as described in the 1125 Service manual. 		

NOTE: Selecting a different drug name in the Drug Library changes the previous concentration and dose rate values to the default values appropriate for the selected drug. The choice of drug names and default values cannot be modified by the user.

The Dose Rate Calculation feature is used to calculate the volumetric rate for a either Channel A or B and is accessed through the menu. The example shown in **Figure 4-7** is for Channel A. However, the dose calculation screen used for channel B is the same except for the title and the soft key text that is used to go to Channel A's dose calculation screen. Once the dose calculation screen has been successfully completed, the dose rate a "Ready to Start" message will appear in the lower, right-hand corner of the display. This message is used to indicate that the user may press the "Start/Stop Channel" control key for the appropriate channel.

The user is free to change the value of any field at any time. When a new infusion rate is entered the dose value will change. When a new dose value, weight, concentration or diluent value is entered the infusion rate changes automatically.

Perform the following steps to select the Dose Rate Calculator:

- a. Press the MENU control key.
- b. Press the DOSE RATE CALC. (A or B) soft key. "Dose Rate Calc. B" message will only be visible if the SidecarTM (Channel B) has been connected.
- c. Observe that the Dose Rate Calculation Menu (See Figure 4-7) will be displayed with the word "Primary" highlighted with post software release 362. The "Dose Value" will be highlighted in software release 362 and earlier (in this case proceed directly to step i).
- d. Use the up and down arrow keys to select Primary, Secondary or Bolus in the post software release 362 version.

NOTE: If the Dose Calculator Drug Library Option is enabled, further choices (*see italics*) are available. If the drug library option is not available, proceed directly to step h. The drug name, patient weight or drug concentration can't be changed while infusing.

- e. Press Enter.
- f. Observe that the word "**Drug**?" is highlighted.
- g. Use the up and down arrow keys to select choices from the included Drug Library.
- h. Press ENTER or the DOSE soft key.
- i. Observe that the Dose value is highlighted. Use the up and down arrow keys to set the desired Dose value. Press ENTER or the DOSE soft key. Observe that the unit is highlighted. Use the up and down arrow keys to choose the correct unit ordered (mg/kg/hr, mg/kg/min, mcg/kg/mi; mcg/kg/hr, mcg/kg, mg/kg, mg/hr, mcg/hr, mg/min, mcg/min).
- j. Press ENTER or the CONC soft key.
- k. Observe that the Conc value is highlighted. Use the up and down arrow control keys to enter the concentration of the medication to be infused. *Press ENTER or the CONC soft key to highlight the Conc units. Use the up and down arrow keys to set the Conc units (mg, mcg).* Press ENTER or the CONC soft key to highlight the base of the concentration if it is greater than 1 mL. Use the up and down arrow keys to select this value.
- 1. Press ENTER or the WEIGHT soft key.
- m. Observe that the Weight value is highlighted.

NOTE: If the patient weight is in pounds, divide that value by 2.2 to get weight in kilograms.

- n. Use the up and down arrow control keys to enter the patient's weight in kg.
- o. Press the appropriate Rate, VTBI, or Time (if displayed) soft key.
- p. Observe that the selected value is highlighted.
- q. Use the up and down arrow control keys to select the value. Press ENTER to complete the selection.
- r. The rate will automatically be calculated and the **Ready to Start** (A) or (B) prompt will appear.
- s. <u>Verify</u> all entered values before starting infusion.

t. Press the START/STOP Channel control key. The Main running screen will display with the correct parameters from the Dose Rate calculation screen with the Dose value displayed between the Rate and VTBI parameters (See Figure 4-8).

·Ⅲ	:55
Primary A 🗦 >	>
>Rate: 1_0 Dose: 0.35 mcg/kg/min	<u>mL</u> hr
> ∨тві: 0	mL
VI: 0.0	mL
≽Bolus	
>Channel B	

Figure 4-8. Primary Screen with Dose Rate Displayed



Figure 4-9. Alarm Volume Adjustment Screen

<u>4.7.2</u> <u>Alarm Volume</u> The MRidiumTM 3850 MRI Infusion Pump allows the clinician to adjust the audio volume for user alerts and alarms (See Figure 4-9). The pump will sound the alarm during adjustment to provide an indication of the current setting of the volume. The volume range is on a visual sliding scale. Perform the following to adjust the Alarm Volume:

- a. Press the MENU control key.
- b. Press the ALARM VOLUME soft key.
- c. Press the up and down arrow control keys to adjust the volume.
- d. Press ENTER.
- e. Press the CANCEL control key to return to the run or programming display screen

NOTE: The Alarm Volume setting in the MENU option of the Pump and Remote Display/ Charger <u>only</u> controls the unit being adjusted. Adjusting the Alarm Volume setting on the Remote Display will <u>not</u> change the Alarm Volume setting of the Pump.

<u>4.7.3</u> <u>KVO Rate.</u> The MRidiumTM 3850 MRI Infusion Pump will default to a preset KVO (Keep Vein Open) rate when the Primary infusion VTBI counts down to 0 mL. This KVO rate may be set to a value between 1-5 mL/hr, according to your hospitals policy, from the Special Feature Setup Menu. Perform the following to adjust the KVO Rate:

- a. Press the MENU control key.
- b. Press the KVO Rate soft key.
- c. Press the soft key corresponding to the Channel you wish to change (if the optional second channel is not present then only Channel A will be available).

NOTE: Always select the appropriate KVO rate for the prescribed fluid therapy.

- d. Use the up and down arrow control keys to adjust limit between 1-5 mL/hr.
- e. Press ENTER.
- f. Press the CANCEL control key to return to the main programming screen.

NOTE: This option may appear on the second Menu page.

<u>4.7.4</u> <u>Occlusion Limits.</u> The MRidiumTM 3850 MRI Infusion Pump will display an Inlet Occlusion alarm when there is an occlusion between the infusion bag and the pump. Patient Occlusion alarm will display when there is an occlusion between the pump and the patient; this alarm has an adjustable pressure limit between 1-10 PSI (6.9 to 68.8 kPa). Occlusion is detected by building of pressure in the I.V. Line downstream of the pump.

NOTE: If the alarm recurs without apparent cause, the Occlusion Pressure Limit may be set too low. Readjust Occlusion Pressure Limit accordingly. Ensure that occlusion pressures chosen are clinically appropriate.

Perform the following to adjust the Occlusion Pressure Limit:

- a. Press the MENU control key.
- b. Press the OCCLUSION LIMIT soft key.
- c. Use the up and down arrow control keys to adjust limit between 1-10 PSI (6.9 to 68.8 kPa).
- d. Press ENTER.

e. Press the CANCEL control key to return to the main programming screen.

NOTE: <u>**Dual Channel Operation:**</u> The KVO Rate and the Occlusion Limit apply to both Channel A and Channel B.

<u>4.7.5</u> <u>Lock Keys.</u> This feature allows the user to prevent an accidental key activation. The Lock Key option will not be displayed unless it has been enabled in the Service Mode.

Perform the following to activate Key Lock:

- a. Press the MENU control key.
- b. Press the LOCK KEYS soft key.
- c. Verify all keys are locked except Menu.
- d. If any other key is pressed a tone will occur to denote that key is locked.

Perform the following to deactivate Key Lock:

- a. Press the MENU control key.
- b. Press the UNLOCK KEYS soft key.
- c. Verify all keys are now unlocked.

This feature is reserved for service personnel to enable, contact Service/Biomedical department for activation (details of this activation are included in the MRidiumTM 3850 MRI Infusion Pump Service Manual, part number 1125). This option may appear on the second Menu page.

<u>4.7.6</u> <u>NEXT MENU key</u> The NEXT MENU will appear if the Remote Display option or the Channel B SideCar option is installed, pressing the NEXT MENU soft key will display additional menu options.

a. **Set Comm Channel:** If the remote control option is installed pressing this soft key will select the Radio Channel selection menu.

WARNING: Upon signal reacquisition between the Pump and the Remote Display, always visually confirm that both Pump and Remote displays change with a selection of the MENU key (this won't affect the Pump operation if it is infusing).

If using multiple Pumps with the same Remote Display/Charger, always program each Pump with its unique radio-link channel (Channels 1 through 6) in the Setup Menu. Additionally, when changing the Remote Display/Charger radio-communication channel, <u>always</u> re-confirm proper communication with the selected Pump. This can be done in many ways, but the easiest way is to select the MENU key on the Remote Display/Charger and visually confirm that the Pump's display matches this change. Pressing the CANCEL key on the Remote Display/ Charger should then return both displays to the original display. Doing this at least once before any Pump programming will acknowledge that proper communication has been established between the Remote Display/Charger and the selected Pump.

<u>4.7.7</u> <u>Radio Channel menu.</u> Press the Channel soft key to highlight the desired channel. Both Remote and Pump must be set to the same channel.



Figure 4-10. Set Comm Channel Displays

<u>4.7.8</u> Exiting the Special Features Menu. To exit the Special Features Menu, press the **MENU** or **CANCEL** control key. The pump will return to the Primary Pump Setup screen where setup of the Primary Pump may be performed.

4.8 Air Bubble Detection and Reset. The pump has an air bubble detector that will detect air bubbles (that are greater than 100 uL) which pass in front of it. When this happens the infusion will stop and the pump will alarm. Once the condition has been corrected the infusion cycle must be manually restarted.

4.9 Data Retention. The pump program settings and option selections are retained in non-volatile memory. If the pump has been turned off for longer than one (1) hour delivery settings are cleared. The History Log data, which stores a comprehensive range of the pumps operational information, is retained indefinitely on a FIFO (First In - First Out) basis. The History Log holds approximately 3,000 to 5,000 entries and alarms. It can be downloaded from the Pump into a computer to review. This is accessed in the Service mode. (Please see Service Manual 1125 for more details).

SECTION 5 ALARMS

5.0 Alarms.

5.1 Introduction.

5.2 User Messages. There are three (3) types of user messages (Alarms, Alerts and User Prompts) displayed on the pump. The following is a description of the user messages:

- a. **Alarm.** Major Pump or Channel related problem.
 - The infusion stops.
 - The RUN/ALARM Lamp illuminates RED and flashes
 - The audible alarm tone sounds
 - An alarm message can appear at top of Main Display
- b. Alert. Indicates a change in the infusion status.
 - The infusion continues to operate
 - The RUN/ALARM Lamp illuminates GREEN and flashes
 - The audible tone sounds
 - An alert message can appear at top of Main Display
- c. **User Prompt.** Informational update, the infusion status has not changed.
 - Generally, some steps were not completed or an incorrect key was pressed.
 - Prompt message can appear at top of Main Display.

NOTE: When using the Pump with both Channel A and Channel B operating, some messages will indicate "Channel A" or "Channel B" to identify which channel is affected. Always verify the appropriate channel is selected before making any changes. If both channels are alarming, both RUN/ALARM indicators will be flashing RED and the alarm message at the top of the Main Display will alternate both messages.

See **Table 5-1** for a listing of all the messages (Alarms, Alerts and User Prompts) that the pump might display. The table is divided into four columns with the first column providing the message which appears on the screen, the second identifies the type of message (Alarm, Alert or User Prompt), the third identifies the cause of the message and the fourth contains the suggested action in response to the message.

- **5.3 Responding to an Alarm.** Perform the following to respond to an alarm:
 - a. Press the Alarm Silence descent control key to silence the sound and re-set the Alarm Condition monitor.
 - b. Resolve the alarm condition (i.e.: clear bubble, change battery, close door, etc.)
 - c. Press the Start/Stop control key for the appropriate channel to continue the infusion.
- 5.4 **Remote Alarms.** All alarms must be resolved at the pump. Alarms are listed in Table 5-1.

WARNING: Always Respond to patient at the pump if an alarm occurs. Do not rely on the remote alarm silence function. Patient injury could occur.

MESSAGE	TYPE	CAUSE	RESOLUTION
Bubble Detected (A) or (B)	Alarm	Air detector has detected an air bubble larger than the 100 uL threshold.	Evaluate air in set. Open Pump Door latch to remove set. Remove air per hospital protocol. Reinstall set. Press START/STOP to resume infusion.
BATTERY DEPLETED	Alert	Battery is too low to operate Pump much longer.	Plug AC Adapter power cord into an AC outlet immediately. Make sure Adapter is plugged into the Pump.
Battery Low	Alert	Battery has 30 minutes or less of charge remaining.	Plug Pump into AC Adapter, and/or AC Adapter power cord into an AC outlet as soon as possible.
Dose Complete (Audible Only)	Alert	A dose delivery has just been completed. Message with an audible beep.	No action necessary. Pump enters KVO.
MAX Rate	Prompt	Calculated rate is outside	Verify and reenter settings.
(Audible Only)		allowable range. An audible beep occurs if key is pressed to attempt to move dose out of range.	
Critical Error xxx	Alarm	Pump Run Test fails, other motor tests, hardware or software conflict.	Remove from use. Contact qualified service personnel.
Close Door (A) or (B)	Alarm	Latch was opened during an infusion.	Check for proper set installation. Close latch. Press START/STOP to resume infusion.
Close Door (A) or (B)	Prompt	Latch is open (prior to starting an infusion). Message occurs with an audible beep that repeats every 10 seconds.	Close latch fully downward.
Patient Occluded (A) or (B)	Alarm	Pressure in I.V. line has exceeded Adjustable Pressure Limit due to elevated resistance in delivery path between pump and patient.	Check administration set for probable cause (kinked tubing, closed stopcock, high resistance catheter, etc.). Press START/STOP key to restart infusion.
Settings Lost	Prompt	Message occurs at power up to indicate the existing operating parameters have been erased due to internal battery failure, and the system has reverted back to initial default parameter settings.	Re-enter patient's infusion settings. If message recurs, refer the pump to qualified service personnel for repair.
DEAD BATTERY	Alarm	Battery too low to operate Pump.	Plug Pump into AC Mains Power. Press ON key and choose "Same Patient" to resume infusion.

 Table 5-1. Alarm, Alert and User Prompt Messages

MESSAGE	TYPE	CAUSE	RESOLUTION
BAD BATTERY	Alert	Battery fault has occurred	Remove from use. Contact qualified service personnel
Inlet Occluded (A) or (B)	Alarm	Flow has been obstructed between fluid container and Pump.	Check administration set for probable cause (kinked tubing, clogged filter, etc.). Press Start/Stop (for appropriate channel) to restart infusion.
Load Set (A) or (B)	Prompt	Pump needs to verify operation (establish pressure baseline and perform motor test).	Open Pump Door until message disappears. Close Door to resume infusion. Always turn Pump on prior to loading set.
Press ENTER	Prompt	Value change confirmation required.	Verify selection and press ENTER.
Pressure Error (A) or (B)	Prompt	Excessive variation in pressure due to motion, flow from other Pumps or blood pressure prevents accurate setting of pressure baseline.	Open and Close door to reset pressure sensors. Press START/STOP key to start infusion.
History Checksum Error	Prompt	Pump detected a memory or power failure. Existing operating parameters have been erased.	Pump has reverted to the default program settings. Press MENU to examine settings, and reenter any needed infusion setting changes.
VTBI = KVO (Audible Only)	Alert	VTBI has counted down to zero. Channel is in KVO mode.	Press the appropriate channel's START/STOP to stop infusion, then press ENTER and the VTBI Soft Key to reenter the new VTBI. Change solution container, if necessary. If therapy is complete, remove tubing from pump to end infusion.
No Communication (Remote Display	Alert	Communication between pump and Remote Display has been interrupted.	Reposition Remote Display or Pump to re-establish communication. Be sure Pump is still ON.
unit only)			Interference from another Radio source may cause problem. Go to an alternate Comm channel on both Pump and Remote Display.
			Otherwise, refer unit to qualified service personnel.
Unload Set	Prompt	Pump needs to reset air in line detector.	Open door. Remove and replace I.V. set. Close door. Always turn Pump ON prior to loading set.
Und	Prompt	Under-Range of Rate or VTBI established settings.	Reset values to be within parameters: Rate: 1.0 to 999 mL/hr VTBI: 0.1 to 9,999 mL
Ovr	Prompt	Over-Range of Rate or VTBI established settings.	Reset values to be within parameters: Rate: 1.0 to 999 mL/hr VTBI: 0.1 to 9,999 mL

MESSAGE	ТҮРЕ	CAUSE	RESOLUTION
Check Door (A) or (B)	Alarm	Possible free flow of I.V. fluid	Be sure door clamp is closed tightly flush with door. Open and reclose door or assure Free Flow Preventer is pulled out to stop position.
Recheck Settings, Press Enter to restart	Prompt	Calculated Dose Rate Calculator values are outside allowable range.	Press the ENTER key. Previous values are replaced with default values. Verify and reenter new settings.
Re-enter settings, press ENTER to continue	Prompt	Selected values are outside allowable range.	Press the ENTER key. Previous values are replaced with default values. Verify and reenter new settings.
Review Primary	Prompt	A selected parameter that affects a Primary infusion calculated value (e.g. Dose, CONC, Weight, etc.) has been modified after setting up an infusion.	Select the Dose Rate Calculator Primary Infusion screen. Infusion can't be started without reviewing this setup screen. Review and verify the new infusion parameter(s) before starting the infusion.
Review Bolus	Prompt	A selected parameter that affects a Bolus infusion calculated value (e.g. Dose, CONC, Weight, etc.) has been modified after setting up an infusion.	Select the Dose Rate Calculator Bolus Infusion screen. Infusion can't be started without reviewing this setup screen. Review and verify the new infusion parameter(s) before starting the infusion.
Review Secondary	Prompt	A selected parameter that affects a Secondary infusion calculated value (e.g. Dose, CONC, Weight, etc.) has been modified after setting up an infusion.	Select the Dose Rate Calculator Secondary Infusion screen. Infusion can't be started without reviewing this setup screen. Review and verify the new infusion parameter(s) before starting the infusion.

SECTION 6 BATTERY OPERATION

6.0 Battery Operation.

6.1 Introduction. The 3850 system includes a smart Battery Pack that self-monitors its performance and status and communicates this information to the Pump. This allows the Pump to monitor the battery parameters in order to maximize the life of the battery, reduce battery-related costs and minimize Pump/Battery down-time.

NOTE: The Battery Pack is sealed such that the batteries can't be replaced. If the Battery Pack fails or becomes unusable, do not attempt to replace the internal batteries. Replace the Battery Pack. Contact Iradimed Corporation for disposal.

6.2 Inserting the Battery Pack. Perform the following to insert the Battery Pack into the pump or Remote Display/Charger:

- a. Grasp the Battery Pack firmly.
- b. Slide the Pack into the battery compartment on the rear of the Unit. Push in firmly until the pack snaps into place. The Battery Pack can be inserted when the Unit is either turned On or Off.
- c. After the Unit is turned On, verify that the Battery Icon on the Main Display has the "X" removed and the Icon is filled or filling during the charge cycle.



Figure 6-1. Battery Installation

6.3 Charging the Battery Pack. After insertion of the Battery Pack, attach the Power Adapter to the Pump and the AC power source. The battery will charge automatically whenever the pump is connected to the AC power source as indicated by a visible Amber LED indicator on the Pump front panel. The Battery Pack can be inserted when the Pump is either turned On or Off.

After the Pump is turned On, verify that the Battery Icon on the Main Display has the "X" removed and the Icon is filled or filling during the charge cycle.

6.4 Removing the Battery Pack. Press in the Battery Pack latch located on the right side of the installed Pack. Grasp the Pack, and pull it outward from the Pump's Battery compartment. Another new or charged battery Pack can now be placed into the Pump. The Battery Pack can be removed when the Pump is either turned On or Off.

WARNING: The Battery Pack is slightly magnetic. Do not remove Battery Pack inside the MRI Scan Room.



Figure 6-2. Battery Removal

6.5 Testing the Battery Pack. The battery level is always evident on the Battery Icon when the Battery Pack is inserted into the Pump, and the Pump is "On". The operating capacity of the Battery Pack can also be tested without the use of the Pump. To examine the capacity level, depress the small button on the rear of the Battery Pack, as indicated on the Battery Pack Label. When pressed the Battery Level LED indicators will momentarily show the capacity (The indicators can range from five (5) L.E.D.s illuminated for fully charged, to one flashing L.E.D. illuminated for fully discharged).



Figure 6-3. Battery Test

6.6 Battery Charge Indicator. The Battery Gauge on the Main Display indicates approximate battery capacity remaining under current operating conditions. It is located in the upper portion of the display and is always visible. Check the remaining battery capacity when starting an infusion. The Battery Gauge updates continuously while infusing.

- a. The battery recharges whenever the Pump and its Power Adapter is plugged into an AC power outlet, or optionally when the Battery Pack is inserted into the Remote Display Unit.
- b. The system includes:
 - A green/amber light that illuminates when Pump/Remote is plugged in to the AC power with the Power Adapter.
 - An amber light that illuminates when Pump is charging the battery, green indicates charging current termination, and the Battery Pack is fully charged.
 - Automatic switchover to battery power if Pump is unplugged or in the event of a power failure.

Note: This automatic switchover is not applicable in the Remote Display Unit, as it cannot operate on battery power.

6.7 Battery Low Indication. A Low Battery alert indicating that battery depletion is imminent, beginning at least 30 minutes prior to a depleted battery alarm. For best results, fully charge, discharge and recharge the battery before putting the Pump into service. Battery run time will be affected by the operating mode, rate, and back pressure.

Should the battery "smart power gauge" detect an over-discharge of one or more internal cells, the pump will provide a "BAD BATTERY" alarm condition. The pump should be turned off and on and recharged immediately to prevent battery damage. Restart/power cycle the pump and select Previous Patient to continue infusion.

CAUTION: If Pump is in storage and not used for more than 30 days, the battery pack should be removed.

6.8 Battery Power Gauge. Located at the top left of the informational display, this gauge is in the shape of a battery and provides a visual indication of the charge status of the battery. When the gauge is "full," then the battery is fully charged; when the gauge is "empty," then the battery is nearly depleted and the pump may not operate unless connected through the MRI Power Supply (pump only) to a hospital grade AC outlet. When the battery is removed, the gauge has an "X" placed inside it. If the Battery Operating time seems short, the battery gauge may require recalibration, to accomplish this, perform the following:

- a. Remove pump from clinical use. Set up pump to operate on battery power.
- b. Operate pump until low power alarm occurs, continue to operate until the "DEAD BATTERY" alarm (capacity <2%) occurs.
- c. Reconnect the pump to the 1120 MRI Power Supply or move the fully depleted battery to the Remote Display/Charger, and allow the pump battery to fully recharge. When in the pump recharging can occur during normal use if the pump is connected to its 1120 MRI Power Supply. The battery should recharge in less than nine (9) hours.
- d. If battery life is still short, refer to qualified service personnel.

6.9 Battery Care and Maintenance:

6.9.1 Introduction. Over discharge, over charge and over current (short-circuit) are all conditions detected by the internal "smart power gauge" of the battery. These conditions all result in the battery shutdown (battery will not deliver power) until placed into charge. Over discharge can cause cells to outgas during subsequent charge cycles. Severe outgassing can result in the battery pack swelling. Replace any battery pack with visible signs of deformation or swelling.

6.9.2 1133 Battery Pack Maintenance Checkout Procedure.

The 1133 Battery Pack should be inspected anytime the pack is removed, or at least every 90 days for the following:

1. <u>Battery Pack communicates with the Pump.</u> This indicated by the Battery Icon being shown on the display of the Pump or Remote Display. Refer to Section 1.3.1 of this manual for Battery Icon description.

2. <u>The Battery Pack holds sufficient operating charge capacity.</u> This is indicated by allowing the Pump to operate on battery power for the recommended minimum time.

3. <u>Battery Pack "Gas Gauge" indicator functions.</u> These functions can be confirmed by performing the checks defined in Sections 6.5 and 6.6 above.

4. <u>Physical Inspection of the Battery Pack case</u>. Remove the Battery Pack from the Pump or Remote Display, and inspect for physical damage or signs of mechanical shock, cracked casing or swollen case.

5. <u>Physical Inspection of the Battery Pack cell(s)</u>. Remove the Battery Pack from the Pump or Remote Display, and confirm that cells inside the Battery Pack are not swelling excessively, as noted by viewing internal cells from the edge of the Battery Pack case. Swollen cells that are greater than 8 mm thick, or cause the Battery Pack's plastic case to bulge indicate a failing battery pack.

Any failure of any of the above inspection criteria will require discontinuance of use and replacement of the 1133 Battery Pack.

6.10 Battery Pack Replacement. If the Battery Pack does not operate for the specified time after a full nine (9) hour charge cycle, replacement of the Battery Pack is recommended.

CAUTION: If placed into storage, to maintain battery life, assure that the battery pack remains charged above 25% (one battery segment is visible) as indicated by the "gas gauge" LED display.

6.11 Battery Pack Related Precautions.

The 1133 Battery Pack contains several lithium-polymer cells and an integral safety circuit. As these cells age, they can expand due to internal gas release, which is anticipated for this type of cell. However, if excessive expansion occurs, this can result in the battery case expanding (swelling), and possibly cause failure of the battery case, cells, or safety circuit. If this is observed, remove the Battery Pack from use and replace it as soon as possible.

The 1133 Battery Pack contains protective circuitry to prevent catastrophic battery failure. If the Battery pack is damaged, this protective circuitry may not prevent battery failure. Remove the Battery Pack from use if the Pack becomes damaged, or the potential for Battery Pack damage is suspected.

Do not use a damaged or swollen 1133 Battery Pack.

Avoid damage to the 1133 Battery Pack by impact, dropping, overheating, or mechanical abuse. Never compress, drop, shock, or strike the 1133 Battery Pack. Never use objects that could puncture the internal battery cells. Any of these actions can cause the battery cells to heat, smoke, or cause catastrophic battery failure, which could result in fire.

Do not attempt to disassemble the 1133 Battery Pack. Damage caused by disassembly or tool use can result in catastrophic battery failure, which could result in fire.

If the 1133 Battery Pack case begins to expand and/or swell, discontinue battery charging and use immediately, and replace the Battery Pack. Continued charging will cause further Battery Pack case expansion, with possible battery case fracture, and potential electrolyte leakage.

If the 1133 Battery Pack becomes damaged, avoid contact with the battery cell electrolyte. If the electrolyte contacts the skin or eyes, seek medical attention immediately.

If the 1133 Battery Pack shows sign of the battery case expanding (swelling), remove the Battery Pack from use and replace it as soon as possible. In extreme conditions, this swelling can cause the 1133 Battery Pack to become jammed or stuck within the 3850 Pump or 3855 Remote Display, and/or cause the Battery Pack plastic case to burst open. If this occurs, do not use tools that could cause damage to the internal battery cells. Refer to the 1125 Service Manual for removal under these conditions.

Under no circumstances should Battery Packs or the internal cells be incinerated as this can cause an explosion.

APPENDIX A SPECIFICATIONS

GENERAL SYSTEM REQUIREMENTS			
General Features	Volumetric Infusion Pump for use in the MRI		
Pump Dive Mechanism	Linear Peristaltic		
Number of Pump Channels	2		
User Event Log	Non-volatile memory retains each operating step and alarm - up to 5000 entries		
ELECTRICAL CHARACTERISTICS			
HI/LO Line Voltage Requirements	100 to 240 VAC +/-10%, 50/60 Hz		
Power Sources Available	Internal battery power with separate AC charger/power supply		
Power Consumption	<15 Volt-Amperes @ 120 VAC nominal at 125 mL/hr (<100 VA maximum during charging)		
Battery Type	Rechargeable Lithium Polymer Pack, 14.8 v at 5.8 Ah		
Battery Capacity	> 12 Hours at 125 mL/hr Rate		
Battery Charge Time	< 9 hours to 95% capacity		
Battery Cycle Life	Typical Life > 200 charge/discharge cycles		
Patient Leakage Current	< 20 uA RMS		
Chassis Leakage Current	< 100 uA RMS; < 300 uA RMS (Single Fault)		
Chassis Ground Impedance to Earth Ground	< 0.1 ohms (with power supply)		
MECHANICAL CHARACTERISTICS			
Dimensions D x W x H	6 x 8 x 9 Inches (15.25 x 20.3 x 22.9 cm)		
Total Weight	10 LBS (4.5 Kg); 11.5 LBS (5.2 Kg) with Battery.		
Operating Temperature Range	+5° to +40° C		
Storage Temperature Range	-40° TO +70° C		
Operating Relative Humidity Range	0% to 80% RH, non-condensing		
Storage Relative Humidity Range	0% to 95% RH, non-condensing		
Pole Clamp Mounting (Pole Diameter) Range	1 inch to 1.5 inch (25 to 38 mm) diameter		

PERFORMANCE CHARACTERISTICS		
Infusion Pump Performance		
Flow Rate Range - 0 to 100 mL/hr	1 to 99.9 mL/hr in 0.1 mL/hr increments	
Flow Rate Range - > 100 mL/hr	100 to 999 mL/hr in 1 mL/hr increments	
Flow Rate Display Range	1.0 to 99.9, 100 to 999 mL/hr	
Flow Rate Accuracy	within 5%	
Primary Volume To Be Infused (VTBI) Range	0.1 to 99.9, 100 to 999 mL	
Secondary Volume To Be Infused (VTBI) Range	0.1 to 99.9, 100 to 999 mL	
Total Volume Infused (VI) Range	0.1 to 99.9, 100 to 9999 mL	
Keep Vein Open (KVO) Rate Range	adjustable, 1 to 5 mL/hr	
Keep Vein Open (KVO) Default Rate	1 mL/hr	
Patient Line (downstream) Back-Pressure Range	+300 to -100 mmHg	
Downstream (proximal) Occlusion Detection Range	1 to 10 PSI (6.9 to 68.8 kPa), user-adjustable	
Occlusion Detection Mechanism	Two separate (upstream & downstream) semiconductor force sensors	
Occlusion Pressure Measurement Range	1 to 10 PSI (6.9 to 68.8 kPa), with 0.2 PSI resolution	
Occlusion Pressure Measurement Accuracy	< 2 PSI (13.8 kPa), or within 10% of setting, whichever is greater.	
Occlusion Detection (no flow) Time	typically <30 sec, dependent on selected Flow Rate, 55 min max. @ 1 mL/Hr	
Occluded I.V. Line bolus volume (25 mL/Hr to 10 PSI occlusion)	0.7 mL max.	
Air-in-Line Detection Method	Ultrasonic bubble detector	
Air-in-Line Detector Threshold(s)	> 100 uL	
Liquid Source height limits	+100 to -50 cm relative to pump center	
Calibration units of measure	mL/hr, mL, Volts, & PSIG	
Audible Alarm Range (at 1 meter)	Minimum: 60 dBA	
	Maximum: >85 dBA	
MRI Performance		
MRI Magnet Compatibility	0.2 to 3.0 Tesla MRI Systems	
MRI Larmor Frequency Interference	No interference at compatible Larmor Frequencies (8.4 through 128 MHz)	
Magnetic Field Limit	10,000 Gauss (1 Tesla magnetic field line) - Minimal ferrous material used inside Pump (< 15 grams); non-magnetic ultrasonic motor used inside Pump	
PRODUCT STANDARDS REQUIREMENTS		
STANDARDS COMPLIANCE: IEC 60601-1	Yes	
STANDARDS COMPLIANCE: IEC 60601-1-2	Yes	
STANDARDS COMPLIANCE: IEC 60601-2-24	Yes	
STANDARDS COMPLIANCE: AAMI ID26	Yes	
STANDARDS COMPLIANCE: UL 60601	Yes	
REMOTE COM	MUNICATIONS	
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FCC Certification	Part 15.247, no license required	
ETS (European) Certification	brETS 300.328, no license required	
Rated RF Output Power	+18 dBm (default), 24 dBm (maximum) Service Selectable.	
Frequency Range	2431.3 to 2474.5 MHz (Iradimed Default) Avoids 802.11 bands 1 and 2	
	France: 2400 to 2450 MHz with Radio programming selection. Service Selectable.	
Number of Channels	85 U.S. Channels. Note: A limited frequency mode is available for France. Service Selectable.	

3851 SIDECAR TM DUAL PUMP DRIVE			
MECHANICAL CHARACTERISTICS			
Dimensions D x W x H	5.5 x 4.5 x 9.5 Inches (14.0 x 11.4 x 24.1 cm)		
Total Weight	6 LBS (2.7 Kg)		
Operating Temperature Range	+5° to +40° C		
Storage Temperature Range	-40° TO +70° C		
Operating Relative Humidity Range	0% to 80% RH, non-condensing		
Storage Relative Humidity Range	0% to 95% RH, non-condensing		
Pole Clamp Mounting (Pole Diameter) Range	1 inch to 1.5 inch (25 to 38 mm) diameter		
ELECTRICAL CH	ARACTERISTICS		
Power Requirements	18 VDC (derived from MRidium Pump)		
Power Consumption	< 19 Volt-Amperes @ 120 VAC nominal		
Chassis Leakage Current	$< 300 \mu A RMS$ (included with MRidium Pump)		
Chassis Ground Impedance to Earth	< 0.1 ohms		
PERFORMANCE C	HARACTERISTICS		
Secondary Pump Performance			
Flow Rate Range - 0 to 100 mL/hr	1 to 99.9 mL/hr in 0.1 mL/hr increments		
Flow Rate Range - > 100 mL/hr	100 to 999 mL/hr in 1 mL/hr increments		
Flow Rate Display Range	1.0 to 99.9, 100 to 999 mL/hr		
Flow Rate Accuracy	within 5%		
Primary Volume To Be Infused (VTBI) Range	0.1 to 99.9, 100 to 999 mL		
Secondary Volume To Be Infused (VTBI) Range	0.1 to 99.9, 100 to 999 mL		
Total Volume Infused (VI) Range	0.1 to 99.9, 100 to 9999 mL		
Keep Vein Open (KVO) Rate Range	adjustable, 1 to 5 mL/hr		
Keep Vein Open (KVO) Default Rate	1 mL/hr		
Patient Line (downstream) Back-Pressure Range	+300 to -100 mmHg		
Downstream (proximal) Occlusion Detection Range	1 to 10 PSI (6.9 to 68.8 kPa), user-adjustable		
Occlusion Detection Mechanism	Two separate (upstream & downstream) semiconductor force sensors		
Occlusion Pressure Measurement Range	1 to 10 PSI (6.9 to 68.8 kPa), with 0.2 PSI resolution		
Occlusion Pressure Measurement Accuracy	< 2 PSI (13.8 kPa), or within 10% of setting, whichever is greater.		
Occlusion Detection (no flow) Time	typically <30 sec, dependent on selected Flow Rate, 55 min max. @ 1 mL/Hr		
Occluded I.V. Line bolus volume (25 mL/Hr to 10 PSI occlusion)	0.7 mL max.		
Air-in-Line Detection Method	Ultrasonic bubble detector		
Air-in-Line Detector Threshold(s)	> 100 uL		
Liquid Source height limits	+100 to -50 cm relative to pump center		

Calibration units of measure	mL/hr, mL, Volts, & PSIG
MRI PERFO	ORMANCE
MRI Magnet Compatibility	Same as Pump (0.2 to 3.0 Tesla MRI Systems)
MRI Larmor Frequency Interference	Same as Pump (No interference at compatible Larmor Frequencies - 8.4 through 128 MHz)
Magnetic Field Limit	Same as Pump (10,000 Gauss (1 Tesla magnetic field line) - Minimal ferrous material used inside unit; non-magnetic ultrasonic motor used inside SideCar Unit)

3855 REMOTE DISPLAY / CONTROL UNIT		
MECHANICAL CH	IARACTERISTICS	
Dimensions D x W x H	8 x 6 x 10 Inches (20.3 x 15.25 x 25.4 cm)	
Total Weight	3.5 LBS (1.6 Kg); 5 LBS (2Kg) with Battery.	
Operating Temperature Range	+5° to +40° C	
Storage Temperature Range	-40° TO +70° C	
Operating Relative Humidity Range	0% to 80% RH, non-condensing	
Storage Relative Humidity Range	0% to 95% RH, non-condensing	
ELECTRICAL CH	ARACTERISTICS	
HI/LO Line Voltage Requirements	100 to 240 VAC +/-10%, 50/60 Hz	
Power Consumption	< 19 Volt-Amperes @ 120 VAC nominal	
	(<100 VA maximum during 1133 Battery Pack charging)	
Spare Battery Charging Bay	For 1133 Rechargeable Lithium Polymer Pack (14.8 v at 6.0 Ah)	
Spare Battery Charge Time	< 9 hours to 95% capacity	
Chassis Leakage Current	< 100 uA RMS; < 300 µA RMS (Single Fault)	
Chassis Ground Impedance to Earth	< 0.1 ohms	
MRI PERF	ORMANCE	
MRI Magnet Compatibility	None specified - 3855 Remote Unit is NOT intended for use inside the Magnet Room	

Drug Name	Label		Adult			Pediatric	
		Primary	Secondary	Bolus	Primary	Secondary	Bolus
Drug?	Dose	1.20 mcg/kg/min	1.20 mcg/kg/min	138.8 mcg/kg/min	1.20 mcg/kg/min	1.20 mcg/kg/min	300 mcg/kg/min
	Conc	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL
	Wt.	60 kg	60 kg	60 kg	15 kg	15 kg	15 kg
	Rate	4.3 mL/hr	4.3 mL/hr	499 mL/hr	1.1 mL/hr	1.1 mL/hr	270 mL/hr
	Time	na	na	na	na	na	na
	VTBI	0.0 mL	0.0 mL	0.0 mL	0.0 mL	0.0 mL	0.0 mL
Adenosine	Dose	140 mcg/kg	140 mcg/kg	140 mcg/kg	140 mcg/kg	140 mcg/kg	140 mcg/kg
	Conc	3 mg/ 1 mL	3 mg/ 1 mL	3 mg/ 1 mL	3 mg/ 1 mL	3 mg/ 1 mL	3 mg/ 1 mL
	Wt.	60 kg	60 kg	60 kg	15 kg	15 kg	15 kg
	Rate	28 mL/hr	28 mL/hr	168 mL/hr	7 mL/hr	7 mL/hr	42 mL/hr
	Time	6 minutes	6 minutes	1 minute	6 minutes	6 minutes	1 minute
	VTBI	2.8 mL	2.8 mL	2.8 mL	0.7 mL	0.7 mL	0.7 mL
							•
Dobutamine	Dose	7.5 mcg/kg/min	7.5 mcg/kg/min	7.5 mcg/kg/min	7.5 mcg/kg/min	7.5 mcg/kg/min	7.5 mcg/kg/min
	Conc	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL	1 mg/ 1 mL
	Wt.	60 kg	60 kg	60 kg	15 kg	15 kg	15 kg
	Rate	27 mL/hr	27 mL/hr	27 mL/hr	6.8 mL/hr	6.8 mL/hr	6.8 mL/hr
	Time	na	na	na	na	na	na
	VTBI	0.0 mL	0.0 mL	0.0 mL	0.0 mL	0.0 mL	0.0 mL
Propofol	Dose	100 mcg/kg/min	100 mcg/kg/min	1.36 mg/kg	180 mcg/kg/min	180 mcg/kg/min	2.5 mg/kg
	Conc	10 mg/ 1 mL	10 mg/ 1 mL	10 mg/ 1 mL	10 mg/ 1 mL	10 mg/ 1 mL	10 mg/ 1 mL
	Wt.	60 kg	60 kg	60 kg	15 kg	15 kg	15 kg
	Rate	36 mL /hr	36 mL/hr	489 mL/hr	16.2 mL/hr	16.2 mL/hr	225 mL/hr
	Time	na	na	1 minute	na	na	1 minute
	VTBI	0.0 mL	0.0 mL	8.2 mL	0.0 mL	0.0 mL	3.8 mL
Dexmedetomidine	Dose	1 mcg/kg/hr	1 mcg/kg/hr	1 mcg/ kg	1 mcg/kg/hr	1 mcg/kg/hr	3 mcg/kg
	Conc	4 mcg/ 1 mL	4 mcg/ 1 mL	4 mcg/ 1 mL	4 mcg/ 1 mL	4 mcg/ 1 mL	4 mcg/ 1 mL
	Wt.	60 kg	60 kg	60 kg	15 kg	15 kg	15 kg
	Rate	15 mL/hr	15 mL/hr	90 mL/hr	3.8 mL/hr	3.8 mL/hr	67.5 mL/hr
	Time	na	na	10 minutes	na	na	10 minutes
	VTBI	0.0 mL	0.0 mL	15 mL	0.0 mL	0.0 mL	11.3 mL

Dose Rate Calculator Default Values

Warning: Default values are provided for initial setup only. Verify all selections prior to beginning the infusion.

US	ER SETTING DEFAULTS	
Characteristic	Specification Range	Initial Start-Up (Factory Default) Value
Primary Flow Rate Range - 0 to 100 mL/ hr	1.0 to 99.9 mL/hr in 0.1 mL/hr increments	1 mL/hr
Primary Flow Rate Range - > 100 mL/hr	100 to 999 mL/hr in 1 mL/hr increments	1 mL/hr
Secondary Flow Rate Range - 0 to 100 mL/hr	1.0 to 99.9 mL/hr in 0.1 mL/hr increments, 100 to 999 mL/hr in 1 mL increments.	1 mL/hr
Primary Volume To Be Infused (VTBI) Range	0.1 to 999 mL	0 mL
Secondary Volume To Be Infused (VTBI) Range	0.1 to 999 mL	0 mL
Total Volume Infused (VI) Range	0.1 to 9999 mL	0 mL
Keep Vein Open (KVO) Rate Range	adjustable, 1 to 5 mL/hr	None, always retains last setting.
Bolus Rate Range	1.0 to 99.9 mL/hr in 0.1 mL/hr increments, 100 TO 999 mL/hr in 1 mL/hr increments	500 mL/hr
Bolus To Be Infused Range	0.1 to 999 mL (max value <= Primary VTBI Setting)	0 mL
Occlusion Pressure Limit Range	1 to 10 PSI in 0.1 PSI increments (6.8 to 68.9 kPa)	None, always retains last setting.
Dose Programming - Patient Weight Range	0.1 to 160 Kg in 0.1 Kg increments	15 = PED 60 = ADULT
Dose Programming - Drug Concentration Range	0.01 to 99.9 mg/mL	Varies, dependent on selected Drug.
Dose Programming CONC - Diluent Range	1 to 999 mL	1 mL
Dose Programming - Dose in mg/Kg/ min Range	0.01 to 9999 mg	Varies, dependent on selected Drug.
Dose Programming - Volume to Be Infused	0.1 to 999 mL	0 mL
Dose Programming - Time Range	1 to 999 min	Varies, dependent on selected Drug.
Alarm Sound Volume	> 65 dBA at 1 meter, Settings= Minimum to Maximum (Graphical Display)	None, always retains last setting.
Drug Library Option	Allow selection of 1 to 5 drugs used in Dose Rate Calculator. When OFF, only one drug (Drug?) is available	Default value is on. (Drug library choices are available)

I.V. ADMINISTRATION SETS	
General Specifications	
IV Set Materials used	PVC Tubing (No Latex or DEHP)
Integral Free Flow Protection	Custom flow preventer incorporated into each set.
	Fluid free-flow can't occur without mechanically
	opening the flow preventer.
Air-Trapping Feature	Air-Trapping "Y" sites used
Access Device Connection	Male Luer Lock fitting provided for connection to
	Access Port
Access Device Connection - needleless port	Yes
IV Set Injection Site Sealing	All sets include self-sealing injection sites
IV Set Maximum Pressure Withstand	> 14 PSI
IV Set Maximum Pull Force Withstand	> 15 Nt (3.3 lbs.), per ISO 8536-4
IV Set Operating Temperature Range	+5' to +40°C
IV Set Storage Temperature Range	-40' TO +52°C
IV Set Storage Shelf Life	4 years
IV Set Infusion-Use Life	6 hours
Recommended Fluid Container Height above	0 to 60 cm (0 to 23.6 inches)
Pump	
1055 Bypass Set	
Primary Set Length	85 inch (215 cm)
Set Inner Diameter	0.108 inch (2.7 mm)
Set Outer Diameter	0.152 inch (3.9 mm)
Priming Volume	12 mL max
1056 Standard Set	
Primary Set Length	118 inch (300 cm)
Set Inner Diameter	0.108 inch (2.7 mm)
Set Outer Diameter	0.152 inch (3.9 mm)
Priming Volume	18 mL max
1057 Syringe Set	
Primary Set Length	85 inch (215 cm)
Set Inner Diameter	0.050 inch (1.3 mm)
Set Outer Diameter	0.092 inch (2.3 mm)
Priming Volume	4 mL max
1058 Extension Set	
Primary Set Length	72 inch (185 cm)
Set Inner Diameter	0.108 inch (2.7 mm)
Set Outer Diameter	0.152 inch (3.9 mm)
Priming Volume	10 mL max

APPENDIX B REPAIR

All repairs on products under warranty must be performed by Iradimed Corporation personnel, or an authorized Iradimed Corporation Service and Repair Center. Unauthorized repairs will void the warranty.

If a pump fails to function properly or requires maintenance, contact Iradimed Corporation Technical Service at 1-407-677-8022 within the U.S., +001-407-677-8022 from outside the U.S.(during normal business hours EST), or by E-mail at techsupport@iradimed.com. Iradimed Corporation Technical Service will advise you of the corrective action required. If you are advised to return the pump to Iradimed Corporation for repair, please do the following:

- a. Obtain a Return Authorization Number. This will ensure proper routing and facilitate timely repair of your pump.
- b. Clean Pump prior to shipment. Do not ship contaminated product to IRadimed Corporation for repair.
- c. Package the pump with adequate protection. If available, use the original carton and packing materials in which the pump was shipped from Iradimed Corporation.
- d. Include a brief description of the problem as well as the name, address and phone number of the person to be contacted for additional information.
- e. Include a purchase order with the pump being returned if it is out of warranty; Iradimed Corporation Technical Services can advise you of your pump's warranty status if need be. Repairs will be made at Iradimed Corporation's current list price for the replacement part(s) plus a reasonable labor charge.
- f. Ship the pump transportation prepaid, to the location specified by your Iradimed Corporation Service Representative with the Return Authorization Number written on the outside of the shipping carton. Repairs will be made, normally, within two (2) weeks and the pump will be returned to you prepaid.

To ensure full reliability, it is recommended that all repairs be made by an Iradimed Corporation Authorized Service and Repair center. For repair at your facility, a competent individual experienced in the repair of pumps can repair the pump only IF it is authorized by Iradimed Corporation Technical Service prior to the repair.

CAUTION: No repair should ever be attempted by anyone not having a complete knowledge of the repair of Iradimed Corporation pumps. Only replace damaged parts with components manufactured or sold by Iradimed Corporation. Contact the Iradimed Corporation Technical Service and Repair Center for service and technical assistance.

APPENDIX C WARRANTY INFORMATION

Iradimed Corporation warrants its major products (i.e. pumps, remote displays, secondary pumps) to be free from defects in materials and workmanship for a period of twelve (12) months from the date of original delivery to the buyer or to buyer's order, provided that same is properly operated under conditions of normal use, and that periodic maintenance and service is performed. A ninety (90) day warranty applies to limited-life parts (e.g.1133 MRI Compatible Battery Pack, 1170 Fiberoptic SpO2 sensor, and Series 1000 infusion sets). A thirty (30) day warranty applies to all parts and accessories not listed above.

This warranty will become null and void if product has been repaired other than by Iradimed Corporation, or its authorized representative, or if the product has been subject to misuse, accident, negligence, or abuse.

Iradimed Corporation's sole obligation under this warranty is limited to repairing a product which has been reported to Iradimed Corporation's Technical Service Center during normal business hours and shipped transportation prepaid. Iradimed Corporation is not liable for any damages including, but not limited to, incidental damages, consequential damages or special damages.

This warranty is in lieu of any other warranties, guarantees or conditions, including merchantability or fitness for a particular purpose. The remedies under this warranty are exclusive and Iradimed Corporation neither assumes nor authorizes anyone to assume for it any other obligation in connection with the sale or repair of its products.

A purchased Maintenance Extension agreement provides for an additional 1, 2, or 3 years of authorized repair for major products. The maintenance extension period will begin at the end of the standard warranty period, and continue until the end of the maintenance extension period purchased. The extended maintenance does not apply to equipment which has been subject to abuse or neglect.

Maintenance Extensions purchased after the standard warranty has expired shall require a physical inspection by Iradimed Corporation prior to purchase of any Maintenance Extension. An additional service fee may also be required to bring the out of warranty product(s) within specifications before any maintenance extension can be activated. (Cost of such inspection and possible repair to the product will be communicated to customer at that time). We reserve the right to refuse the sale of Maintenance Extension to any Product.

Iradimed Corporation warrants any such product subject to a Maintenance Extension agreement shall, other than its expendable parts, provided that same is properly operated under conditions of normal use, and that periodic maintenance and service is performed; be repaired by Iradimed and restored to full operational specification as where applicable at the time of original manufacture. Any Maintenance Extension will become null and void if product has been repaired other than by Iradimed Corporation, or its authorized representative, or if the product has been subject to misuse, accident, negligence or abuse.

Should a unit perform outside of Iradimed specifications and cannot be corrected by on site technicians with instruction and support from Iradimed and unit must be returned to Iradimed for repair, a loaner unit, if available, may be provided.

IRADIMED CORPORATION PRODUCTS CONTAIN PROPRIETARY COPY WRITTEN MATERIAL; ALL RIGHTS ARE RESERVED BY IRADIMED CORPORATION

APPENDIX D

MANUFACTURERS TECHNICAL DECLARATION

EMC Information Tables as required by EN 60601-1-2:2000 Clause 6

In accordance with EN 60601-1-2:2000 Medical Electrical Equipment – Part 1-2: General requirements for safety – Collateral standard: Electromagnetic Compatibility – Requirements and tests

- 1. "Medical Electrical Equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in the Accompanying Documents" (the following tables).
- 2. "Portable and Mobile RF Communications Equipment can affect Medical Electrical Equipment".
- 3. "The equipment or system should not be used adjacent to or stacked with other equipment and that if adjacent or stacked use is necessary, the equipment or system should be observed to verify normal operation in the configuration in which it is used".

The following tables (as required in EN 60601-1-2:2000) provide information regarding the Electromagnetic Compatibility (EMC) of this product and its accessories.

Table 201—Guidance and manufacturer's declaration— electromagnetic emissions—for all EQUIPMENT and SYSTEMS

Guidance	and manufacturer's de	eclaration—electromagnetic emissions
The MRidium 3850	System is intended for	use in the electromagnetic environment specified below. The
customer or the use	er of the MRidium 3850	System should assure that it is used in such an environment.
Emissions Test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1	The MRidium 3850 System must emit electromagnetic energy in order to perform its intended function (remote communications within an specific band for WLAN; i.e. 2.431 to 2.474 GHz). Nearby electronic equipment may be affected.
RF emissions CISPR 11	Class B	The MRidium 3850 System is suitable for use in all establishments other than domestic and those directly
Harmonic emissions IEC 61000-3-2	Not applicable	connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/flicker emissions IEC 61000-3-3	Not applicable	

MRIdium 3850 System

Table 202—Guidance and manufacturer's declaration— electromagnetic immunity—for all EQUIPMENT and SYSTEMS

The MRidium 3850 System is intended for use in the electromagnetic environment customer or the user of the MRidium 3850 System should assure that it is used in such an environment. Immunity test IEC 60601 test level Compliance level Electromagnetic environment - guidance level Electrostatic discharge (ESD) ±6 kV contact ±6 kV contact ±8 kV air Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %. Electrical fast ±2 kV for power supply lines ±2 kV for power supply lines Mains power quality should be that of a typical commercial or hospital environment. Surge ±1 kV for input/output lines ±1 kV for input/output lines ±1 kV differential mode Mains power quality should be that of a typical commercial or hospital environment. Voltage dips, short ±5 % UT (>95 % dip in UT) for 0.5 cycle <5 % UT (>95 % dip in UT) for 0.5 cycle Mains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power mains interruptions, it is recommended that the MRidium 3850 System be powered from an uninterruptible power supply in UT) for 5 cycles KD % UT (30 % dip in UT) for 5 sec <5 % UT(>95 % dip in UT) for 5 sec <5 % UT(>95 % dip in UT) for 5 sec Power frequency (50/60 Hz) magnetic field 3 A/m 3 A/m Power frequency magnetic fields should be at levels cha		Guidance and ma	nufacturer's declaratio	on—electromagnetic immunity
Immunity test IEC 60601 test level Compliance level Electromagnetic environment - guidance Electrostatic discharge (ESD) ±6 kV contact ±6 kV contact ±6 kV contact Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %. Electrical fast transient/burst ±2 kV for power supply lines ±2 kV for power supply lines ±2 kV for power supply lines ±2 kV for input/output lines Mains power quality should be that of a typical commercial or hospital environment. Surge ±1 kV differential mode ±1 kV differential mode ±1 kV differential mode Mains power quality should be that of a typical commercial or hospital environment. Voltage dips, short interruptions, and voltage variations on power supply input lines <5% UT (>95 % UT (>95 % UT (>95 % dip in UT) for 5 cycles 70 % UT (30 % dip in UT) for 25 cycles 70 % UT (30 % dip in UT) for 5 cycles Mains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power continued operation during power mains interruptions, it is recommended that the MRidium 3850 System requires continued operation during power and in uninterruptions, it is recommended that the MRidium 3850 System supply or a battery. Power frequency (50/60 Hz) magnetic field 3 A/m Power frequency magnetic fields s	The MRidium 3850 System	is intended for use	in the electromagnetic e	nvironment customer or the user of the MRidium
Immunity testIEC 60601 test levelCompliance levelElectromagnetic environment - guidanceElectrostatic discharge (ESD) IEC 61000-4-2±6 kV contact ±8 kV airFloors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.Electrical fast transient/burst±2 kV for power supply lines±2 kV for power supply linesMains power quality should be that of a typical commercial or hospital environment.Surge IEC 61000-4-5±1 kV differential mode±1 kV differential mode±1 kV differential modeMains power quality should be that of a typical commercial or hospital environment.Voltage dips, short interruptions, and voltage variations on power supply input lines<5% UT (>95 % UT (>95 % UT OW UT (30 % dip in UT) for 5 cycles<5% UT (>95 % UT (30 % dip in UT) for 5 cyclesMains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power romins interruptions, it is recommended that the MRidium 3850 System requires continued operation during power romins interruptions, it is recommended that the MRidium 3850 System requires continued operation during power anios interruptions, it is recommended that the MRidium 3850 System requires continued operation during power anios interruptions, it is recommended that the MRidium 3850 System requires continued operation during power anios interruptions, it is recommended that the MRidium 3850 System is recommended that the MRidium 3850 System is recommended that the MRidium 3850 System is power of from an uninterruptible power system supply or a battery.	3850 System should assure	e that it is used in su	ich an environment.	
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Electrostatic discharge (ESD) ±6 kV contact ±6 kV contact Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %. Electrical fast ±2 kV for power supply lines ±2 kV for power supply lines Mains power quality should be that of a typical commercial or hospital environment. Surge ±1 kV differential mode ±1 kV differential mode ±1 kV differential mode Mains power quality should be that of a typical commercial or hospital environment. Voltage dips, short interruptions, and voltage uput lines <5 % UT (>95 % dip in dip in UT) for 0.5 <5 % UT (>95 % dip in UT) for 5 cycles Mains power quality should be that of a typical commercial or hospital environment. ELE 6 1000-4-11 40 % UT (60 % dip in UT) for 5 cycles <5 % UT (>95 % dip in UT) for 5 cycles Mains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power mains interruptions, it is recommended that the MRidium 3850 System be powered from an uninterruptible power supply or a battery. Power frequency (50/60 Hz) magnetic field Hz magnetic field 3 A/m 3 A/m Power frequency magnetic fields should be at typical commercial or hospital environment.		level		
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IEC 61000-4-2 ±8 kV air ±8 kV air the relative numidity should be at least 30 %. IEC 61000-4-2 ±2 kV for power supply lines ±2 kV for power supply lines Mains power quality should be that of a typical commercial or hospital environment. IEC 61000-4-4 ±1 kV for input/output lines ±1 kV for input/output lines ±1 kV differential mode Mains power quality should be that of a typical commercial or hospital environment. Surge ±1 kV differential mode ±1 kV differential mode ±1 kV common mode Mains power quality should be that of a typical commercial or hospital environment. Voltage dips, short interruptions, and voltage variations on power supply input lines <5 % UT (>95 % dip in dip in UT) for 0.5 cycles Wair (60 % dip in UT) for 5 cycle Mains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power continued operation during power mains interruptions, it is recommended that the MRidium 3850 System be powered from an uninterruptible power supply or a battery. Power frequency (50/60 Hz) magnetic field Hz) magnetic field 3 A/m Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	(ESD)		.0.1)/	tile. If floors are covered with synthetic material,
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IEC 61000-4-4±1 kV for input/output lines±1 kV for input/output lines±1 kV for input/output linesSurge±1 kV differential mode±1 kV differential mode ±2 kV common modeMains power quality should be that of a typical commercial or hospital environment.IEC 61000-4-5±2 kV common mode±2 kV common modeMains power quality should be that of a typical commercial or hospital environment.Voltage dips, short interruptions, and voltage variations on power supply input lines<5 % UT (>95 % UT (>95 % cycles<5 % UT (>95 % dip in UT) for 0.5 cycleMains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power continued operation during power mains interruptions, it is recommended that the MRidium 3850 System be powered from an uninterruptible power supply or a battery.Power frequency (50/60 Hz) magnetic field IEC 61000-4-83 A/m3 A/mPower frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	transient/burst	supply lines	lines	commercial or hospital environment.
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IEC 61000-4-5mode ±2 kV common mode±2 kV common modecommercial or hospital environment.Voltage dips, short interruptions, and voltage variations on power supply input lines<5 % UT (>95 % dip in dip in UT) for 0.5 cycleMains power quality should be that of a typical commercial or hospital environment. If the user of the MRidium 3850 System requires continued operation during power continued operation during power mains interruptions, it is recommended that the MRidium 3850 System be powered from an uninterruptible power supply or a battery.Power frequency (50/60 H2) magnetic field IEC 61000-4-83 A/m3 A/mPower frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	Surge	±1 kV differential	±1 kV differential mode	Mains power quality should be that of a typical
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Hz) magnetic field IEC 61000-4-8	Power frequency (50/60	3 A/m	3 A/m	Power frequency magnetic fields should be at
IEC 61000-4-8 typical commercial or hospital environment.	Hz) magnetic field	57.VIII	0,011	levels characteristic of a typical location in a
	IEC 61000-4-8			typical commercial or hospital environment.

NOTE—UT is the a.c. mains voltage prior to application of the test level.

The MRidium	3850 System is intend	ed for use in the electro	pmagnetic environment customer or the user of the MRidium
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
			Portable and mobile RF communications equipment should be used no closer to any part of the MRidium 3850 System, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance
Conducted RF	3 Vrms 150 kHz to 80 MHz outside ISM bandsª	3 V	d = 1.17 √ P
IEC 61000-4-6	10 Vrms 150 kHz to 80 MHz outside ISM bandsª	10 V	d = 1.20 √ P
Radiated RF	10 Vrms 80 MHz to 2.5 GHz	10 V/m	d = 1.20 √ P 80 MHz to 800 MHz
IEC 61000-4-3			d = 2.33 √ P 800 MHz to 2.5 GHz
			where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m).b
			Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, c should be less than the compliance level in each frequency range.d
			Interference may occur in the vicinity of equipment marked with the following symbol:
			IEC 60417, No. 417-IEC5140, "Source of Non-Ionizing Radiation" Symbol

Table 203—Guidance and manufacturer's declaration— electromagnetic immunity—for LIFE-SUPPORTING EQUIPMENT and SYSTEMS

NOTE 2—These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, obje

and people. a The ISM (industrial, scientific, and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz

b The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz are intended to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in these frequency ranges.

c Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast, and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the MRidium 3850 System is used exceeds the applicable RF compliance level above, the MRidium 3850 System should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the MRidium 3850 System.

Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [V1] V/m

MRIdium 3850 System

Table 205—Recommended separation distances between portable and mobile RF communications equipment and the EQUIPMENT or SYSTEM—for LIFE-SUPPORTING EQUIPMENT and SYSTEMS

Recommended separation distances between portable and mobile RF communications equipment and the MRidium 3850 System

The MRidium 3850 System is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the MRidium 3850 System can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the MRidium 3850 System as recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separa	ation distance accordir	ng to frequency of trans	smitter
output power of transmitter	150 kHz to 80 MHz outside ISM bands	150 kHz to 80 MHz in ISM bands	80 MHz to 800 MHz	800 MHz to 2.5 GHz
••	d = 1.17 √ P	d = 1.2 √ P	d = 1.2 √ P	d = 2.33 √ P
0.01	0.12	0.12	0.12	0.23
0.1	0.38	0.38	0.38	0.77
1	1.17	1.2	1.17	2.33
10	3.8	3.8	3.8	7.67
100	11.67	12	11.67	23.33

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be determined using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1—At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE 2—The ISM (industrial, scientific, and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

NOTE 3—An additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 4—These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Application of Council Directive(s) R&TTE Directive 1999/5/EC Regulation(s) to which Conformity is Declared <u>EN 300 328, IEC 60601-1, IEC 60601-1-2</u> Standard(s) to which Conformity is Declared <u>EN 300 328, IEC 60601-1, IEC 60601-1-2</u> Manufacturer's Name Iradimed Corporation Manufacturer's Address 7457, Aloma Ave., Winter Park, FL 32792 USA Importer's Name Refer to accompanying Packing Slip Importer's Address Refer to accompanying Packing Slip Cype of Equipment MRIdium 3850 Infusion Pump System and accessories Model No(s). 3850 Series and 3855 Series Serial No(s). Refer to accompanying Packing Slip Year of Manufacture Refer to accompanying Packing Slip Zertification Method(s) Self-certified		DECLARATION OF CONFORMITY
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Model No(s). 3850 Series and 3855 Series Serial No(s). Refer to accompanying Packing Slip Year of Manufacture Refer to accompanying Packing Slip Certification Method(s) Self-certified	Type of Equipment	MRIdium 3850 Infusion Pump System and accessories
Serial No(s). Refer to accompanying Packing Slip Year of Manufacture Refer to accompanying Packing Slip Certification Method(s) Self-certified	Model No(s).	3850 Series and 3855 Series
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	Place <u>Winter Park, FL</u>	Junii Carly
Place Winter Park, FL	Date <u>April 10, 2007</u>	(Signature) V Francis Casey (Full Name) Regulatory Affairs

Declaration of Conformity with Regard to the EU Directive 1999/5/EC (R&TTE Directive)

Scope

The information in this document is applicable to the MRidium 3850 System using the 3850R MRI Infusion Pump with 2.4 GHz FHSS Transceiver, and 3855 Remote Display Controller with the 2.4 GHz FHSS Transceiver, and other related products as they come to market.

The equipment operates in the bands (or portions of) 2400 to 2483.5 MHz. National regulations may require that operations be limited to a portion of the frequency ranges identified above. See the National Restrictions section for full details.

Czech (CS) Česky	Toto zařízení je v souladu se základními požadavky a ostatními odpovídajícími ustanoveními směrnice 1999/5/ES
Danish (DA) Dansk	Dette udstyr er i overensstemmelse med de væsentlige krav og andre relevante bestemmelser i Direktiv 1999/5/EF.
Dutch (NL) Nederlands	Dit apparaat voldoet aan de essentiele eisen en andere van toepassing zijnde bepalingen van de Richtlijn 1999/5/EG
Estonian (ET) Eesti	See seade vastab direktiivi 1999/5/EÜ olulistele nõuetele ja teistele asjakohastele sätetele.
English (EN)	This equipment is in compliance with the essential requirements and other relevant provisions of Directive 1999/5/EC.
German (DE) Deutsch	Dieses Gerät entspricht den grundlegenden Anforderungen und den weiteren entsprechenden Vorgaben der Richtlinie 1999/5/EG
Greek (EL) Ελληνική	Αυτός ο εξοπλισμός είναι σε συμμόρφωση με τις ουσιώδεις απαιτήσεις και άλλες σχετικές διατάξεις της Οδηγίας 1999/5/ΕΚ
Finnish (FI) Suomi	Tämä laite täyttää direktiivin 1999/5/EY olennaiset vaatimukset ja on siinä asetettujen muiden laitetta koskevien määräysten mukainen.
French (FR) Français	Cet appareil est conforme aux exigencies essentielles et aux autres dispositions pertinentes de la Directive 1999/5/CE
Hungarian (HU) Margyar	Ez a készülék teljesíti az alapvető követelményeket és más 1999/5/EK irányelvben meghatározott vonatkozó rendelkezéseket.
lcelandic (IS) Íslenska	Þetta tæki er samkvæmt grunnkröfum og öðrum viðeigandi ákvæðum Tilskipunar 1999/5/EB
Italian (IT) Italiano	Questo apparato è conforme ai requisiti essenziali ed agli altri principi sanciti dalla Direttiva 1999/5/CE
Latvian (LV) Latviski	Šī iekārta atbilst Direktīvas 1999/5/EK būtiskajām prasībām un citiem ar to saistītajiem noteikumiem.
Lithuanian (LT) Lietuvių	Šis įrenginys tenkina 1999/5/EB Direktyvos esminius reikalavimus ir kitas šios direktyvos nuostatas.

Declaration of Conformity

Maltese (MT) Malti	Dan I-apparat huwa konformi mal-htigiet essenzjali u I-provedimenti I-ohra rilevanti tad- Direttiva 1999/5/KE
Norwegian (NO) Norsk	Dette utstyret er i samsvar med de grunnleggende krav og andre relevante bestemmelser i EU-direktiv.1999/5/EC
Polish (PL) Polski	Urządzenie jest zgodne z ogólnymi wymaganiami oraz szczególnymi warunkami określonymi Dyrektywą UE: 1999/5/WE
Portuguese (PT) Português	Este equipamento está em conformidade com os requisitos essenciais e outras provisos relevantes da Directiva 1999/5/CE

Applicable Standards

The following standards were applied during the assessment of the product against the requirements of the Directive 1999/5/EC:

- Radio: EN 300 328
- EMC: EN 60601-1-2
- Safety: EN 60601-1-1 and EN 60601-2-24

CE Marking

For the 2.4 GHz wireless products such as the MRidium 3850 System using the 3850R MRI Infusion Pump with 2.4 GHz FHSS Transceiver, and 3855 Remote Display Controller with the 2.4 GHz FHSS Transceiver, the following CE mark and class-2 identifier are added to the equipment:

CE (D)

National Restrictions

In the majority of the EU and other European countries, the 2.4 GHz band has been made available for the use of ISM (Industrial, Scientific, and Medical) communication. Table 1 provides an overview of the regulatory requirements in general that are applicable for the 2.4 GHz band.

Later in this section you will find an overview of countries in which additional restrictions or requirements or both are applicable.

The requirements for any country may evolve. Iradimed Corporation recommends that you check with local authorities for the latest status of their national regulations for 2.4 GHz ISM (Industrial, Scientific, and Medical) communication.

Frequency Band:	2400 - 2483.5 MHz

Max Power Level (EIRP): 100 mW

Indoor and Outdoor Use: Yes

The following sections identify countries having additional requirements or restrictions than those listed above:

France

For 2.4 GHz, the output power is restricted to 10 mW EIRP when the product is used outdoors in the band 2454 - 2483.5 MHz. There are no restrictions when used in other parts of the 2.4 GHz band. For use in France, channel selection is limited to 2400 to 2450 MHz. Check http://www.arcep.fr/ for more details.

Italy

This product meets the National Radio Interference specifications and the requirements in the National Frequency Allocation Table for Italy. Operating wireless equipment requires a "general authorization" unless it is operated within the boundaries of the owner's property. Please check with <u>http://comumicazioni.it/it</u> for more information.

Questo prodotto è conforme alla specifiche di Interfaccia Radio Nazionali e rispetta il Piano Nazionale di ripartizione delle frequenze in Italia. Se non viene installato all'interno del proprio fondo, l'utilizzo di prodotti Wireless LAN richiede una "Autorizzazione Generale". Consultare http://www.comunicazioni.it/it per maggiori dettagli.

Latvia

Any outdoor usage of the 2.4 GHz band requires an authorization form the Electronic Communications Office. Please check <u>http://www.esd.lv</u> for more information.

Antennas

The 2.4 GHz wireless ISM (Industrial, Scientific, and Medical) communication products described in this document have dedicated antennas which should not be removed or replaced with any antenna that is not supplied directly from Iradimed Corporation or its representatives.

Operating Frequency

The operating frequency is pre-set at the factory during manufacturing. Manual setting of the operating frequency can only be done by qualified service personnel with a service

password. Instructions are provided in the Service documentation regarding the appropriate choices and settings for specific geographic areas. It is important that the operating frequency is correctly configured to meet the local regulations.

Changing Output Power

The output power is pre-set to 63 mW at the factory during manufacturing. Manual setting of the output power above 100 mW can only be done by qualified service personnel with a service password. The only available ouput power choices are Level 1 (10 mW), Level 2 (63 mW – factory default), or Level 3 (250 mW). Instructions are provided in the Service documentation regarding the appropriate choices and settings for specific geographic areas. It is important that the output power is correctly configured to meet the local regulations.

Note If you still have questions regarding the compliance of these products, or you cannot find the information you are looking for on the Iradimed Corporation website, http://iradimed.com, please send an email request to Iradimed Corporation at:

techsupport@iradimed.com

APPENDIX E TROUBLESHOOTING

Problem	Possible Cause	Solution
Pump won't turn on.	Unit not turned on.	Turn on unit by pressing 1 (On) control key.
	No AC power/battery depleted.	Plug power cord into pump and AC power source or replace battery pack with charged battery.
	Blown fuse(s).	Refer to qualified service personnel.
	AC power source has low voltage.	Switch AC power cord to power receptacle with sufficient voltage.
No Fluid Flow.	Administration Set is not connected to fluid source.	Attach Administration Set to fluid source.
	Set is not mounted in pump.Open pump door and re-load Administration Set.	
	Administration Set does not mount onto pump.	Select appropriate MRidium TM 3850 Administration Set for pump.
	Patient line is kinked.	Straighten kinked line. Inspect for damage or leaks, and replace if necessary.
	Roller Clamp is closed.	Open Roller or slide clamp.
	Damaged Set or line is leaking.	Replace damaged Administration Set.
Pump won't start.	Door Open.	Reseat door, close completely.
	Set is not properly mounted in pump.	Open pump door and re-load Administration Set.
	Pump program has not been entered.	Enter Pump settings and initiate Pump by pressing START/STOP for appropriate channel.

Problem	Possible Cause	Solution	
Eluid Lasting from	Fluid side of Set is not properly connected.	Inspect for damage or leaks, and replace if necessary.	
Administration Set.	Damaged Set is leaking.	Examine Set to locate leak. Disconnect Set from the patient. Replace and resume infusion.	
	Battery not properly charged.	Plug pump power cord into AC power source until battery is fully charged.	
Unit won't operate on battery.	Aged battery won't hold charge.	Plug pump power cord into AC power source until battery is fully charged. If battery won't charge, replace battery pack.	
	Battery not properly charged.	Plug pump power cord into AC power source until battery is fully charged.	
Battery operation is too short.	Aged battery won't hold charge.	Plug pump power cord into AC power source until battery is fully charged. If battery won't charge, replace battery pack and charge again. If problem persists, refer to qualified service personnel.	
Battery is hard to remove from Pump.	Battery cell failure due to over discharge or individual cell failure.	Remove/replace battery in pump, using appropriate tools as needed. Refer damaged battery to qualified service personnel.	
No audible alarm heard.	Occlusion Pressure limits not set properly.	Adjust limits to clinically appropriate level.	
	Alarm volume set too low for the use environment.	Adjust alarm volume for the intended use environment.	
	Faulty alarm speaker.	Refer to qualified service personnel.	
Continuous audible alarm heard.	Faulty pump hardware.	Refer to qualified service personnel.	
Fluid delivery seems inaccurate.	Pump not calibrated.	Refer to qualified service personnel.	
Keys won't function	Key not pressed firmly.	Repeat key press pushing in firmly.	
Reys wont function.	Faulty key panel.	Refer to qualified service personnel.	

Problem	Possible Cause	Solution	
Pump settings are restored to default.	Pump turned off for longer than one (1) hour between uses.	Train appropriate personnel that turning off the pump for more than one (1) hour resets pump settings.	
	Pump settings changed accidentally.	Double check key sequence for feature chosen for correct key sequence. Was "New Patient" chosen?	
	History Checksum Error, internal battery problem.	Refer to qualified service personnel.	
History Log data are lost accidentally.	Pump memory is damaged or internal backup battery is low.	Refer to qualified service personnel. Memory can only be reset by manufacturer.	
Infusion prematurely stopped.	Occlusion overpressure, inlet occlusion or patient occlusion.	Examine Set to locate occlusion. Disconnect Set from the patient to clear occlusion. Re-prime line and resume infusion.	
	Bubble Detected alarm occurred.	Examine Set to locate air bubble. Disconnect Set from the patient. Purge Air in Line. Re-prime line and resume infusion.	
	Faulty pump hardware.	Refer to qualified service personnel.	
Unit alarms after each infusion start.	Occlusion pressure alarm limit is set too low.	Examine patient infusion site for source of occlusion or infiltration. Re- adjust alarm limits as described in Paragraph 4.7.4	
	Bubble Detected alarm occurred.	Examine Set to locate air bubble. Disconnect Set from the patient. Purge Air in Line. Re-prime line and resume infusion.	
Pump clock is not correct.	Clock not properly adjusted to local time after receipt.	Adjust clock time as described in service manual.	
	Clock battery has become depleted.	Refer to qualified service personnel.	
	Summer/Winter time change.	Adjust clock as described in service manual.	

Problem	Possible Cause	Solution
Infusion stops.	Air is present in the Administration Set.	Examine Set to locate air bubble. Disconnect Set from the patient. Purge Air in Line. Re-prime line and resume infusion.
	Administration Set Roller clamp is closed.	Open Roller Clamp on Administration Set.
	Administration Set improperly loaded on pump drive.	Open pump door, remove Set and reload Set following instructions.
	Kinked Set causes Occlusion Alarm.	Straighten Set and re-start infusion.
	Patient Access Device is blocked.	Change Patient Access Device.
Inlet Occlusion Alarm occurs when using 1057 Syringe IV Set	1057 IV set air vent is not open, which prevents normal fluid flow.	Open the white cap of the 1057 IV set air vent to allow air flow into the syringe.
	Occlusion Pressure Alarm limit is set too low for the smaller diameter IV tubing.	Select a higher Occlusion Pressure Alarm limit to allow normal flow. Alarm limits may need to be set between 8 and 10 PSI, depending on fluid temperature and viscosity.
Recurring "Check Door" or "Close Door" messages occur when pump door is closed.	The Door Clamp rubber pin is bent or damaged.	The Door Clamp rubber pin needs replacement. Refer to qualified service personnel.

Problem	Possible Cause	Solution
Remote Display will not communicate with selected pump	Pump is not within range.	Reposition pump to establish communication. Verify pump is within 90 ft (30 m) of Remote Display.
	Pump is not operational.	Verify pump power is on and radio antenna is securely attached to both Pump and Remote Display.
	Pump and Remote Display are not set at the same channel.	Verify in the Set Comm Channel menu option that both Pump and Remote Display are set to the same comm channel. If not, change the Remote Display to the Pump's comm channel.
	Local radio interference prevents communication.	Select alternate comm channels for both Pump and Remote Display. However, follow the installation procedure (Remote Display Installation in Section 2) to avoid the possibility of setting 2 Pumps on the same channel.
	MRI Magnet Room shielding reduces radio signal strength to prevent communication.	Refer to qualified service personnel. Radio transmission power can be increased (within limits) by service personnel to accommodate most MRI magnet rooms. Contact Iradimed Corporation for additional information.

APPENDIX F ACCESSORIES

Infusion Sets

1055-50 Bypass Infusion Set (carton of 50)
1056-50 Infusion Set (carton of 50)
1057-50 Syringe Adapter Set (carton of 50)
1058-50 Extension Set (carton of 50)

1119 - MRidiumTM MRI IV Pole (Non-Magnetic (non-ferrous) I.V. Pole, 1.5 in.dia.)

- 1120 3850 Pump MRI Power Supply
- 1121 AC Power Adapter Power Cord, 3 ft. (length), medical grade
- 1125 MRidium Service Manual CD
- 1122 AC Adapter/To Pump Interconnect Cable, 10 ft. (length)
- 1127 Quick Reference Hang Tags
- 1128 AC Power Cord, 3.05 m (length), medical grade
- 1133 MRI-Compatible Battery Pack
- 3851 Channel B Pump Drive Assembly (Side-mount)
- 3855 Remote Display/Battery Charger
- LB2025 Channel ID Marker Sheet
- AM01 MRidium[™] 3850 Pump System Software Update Card

Note: The 3850R designates a 3850 Pump with the optional 2.4 GHz wireless link for communication with the 3855 Remote Display.

WARNING: Use only listed accessories with the MRidium[™] 3850 MR Pump.

APPENDIX G TRUMPET AND START UP CURVES

In this Pump, as with all infusion systems, the actions of the pumping mechanism can cause short term fluctuations in rate accuracy. The graphs below (Figure G-1 through G-14) show typical performance of the system with its specific infusion sets.

For those not familiar with trumpet curves, the "0" line in each Figure represents the set flow for the pump and quantities above this line represent the percentage by which the delivered flow exceeds the set flow rate. Quantities below the line represent a flow rate that delivers less than the set flow rate. The heavy, dark solid line represents the average flow rate error for the entire test interval, typically less than 3%.

The maximum positive and negative mean flow errors are shown at 2, 5, 11, 19, and 31 minute averages, and the trumpet curves interpolate the data between these points. The longer the time interval is, the narrower the error range will be. For example, an interval of 15 minute will show a narrower error range than an interval of 3 minute. This is because, at longer observation intervals, fluctuations — even large ones — occurring for only a few minutes represent a comparatively small portion of the data points being analyzed. Conversely, for a shorter interval, any fluctuation will have greater weight.

Because of this weighting, the plotted curve takes on its trumpet shape, with the bell of the trumpet widening at the shorter observation periods. For further information refer to <u>Health</u> <u>Devices</u> Vol. 27 Nos. 4-5 (April-May 1998).

As detailed in the product standards listed below, trumpet curves are named for their characteristic shape. They display discrete accuracy data averaged over particular time periods or "observation windows", not continuous data versus operating time. Over long observation windows, short-term fluctuations have little effect on accuracy, as represented by the flat part of the curve. Variations in system accuracy and start up characteristics over various observation windows can be of interest when certain drugs are being delivered. Trumpet curves are all derived from the second hour of the data collection period.

FLOW CHARACTERISTICS UNDER VARYING DELIVERY CONDITIONS

Effects of Pressure Variations

Under conditions of +300 mmHg pressure, the MRidiumTM 3850 Infusion System exhibits a long-term accuracy offset of approximately -1.0% from mean values. Under conditions of -100 mmHg pressure, the MRidiumTM 3850 Infusion System exhibits a long-term accuracy offset of approximately +3.0% from mean values. No significant change in short–term variations result under these pressure conditions.

Effects of Negative Solution Container Heights

With a negative head height of -0.5 meters, the MRidiumTM 3850 Infusion System exhibits a long–term accuracy offset of approximately -2.0% from mean values. No significant change in short–term variations result under negative head height conditions.

Effects of Rate

No significant change in short-term variations result with rates of 1.0 ml/hr or above.

NOTE: Tests were conducted in accordance with IEC 60601–2–24, "Particular requirements for safety of infusion pumps and controllers" and AAMI ID26–1998 "Medical electrical equipment - Part 2: Particular requirements for the safety of infusion pumps and controllers." MRidium 1056 Infusion Sets were used during this testing.

Occlusion Pressure Tests

NOTE: 2 PSI = 13.8 kPa, 10 PSI = 68.8 kPa

Occlusion Pressure Test Conditions	Occlusion Pressure Test Results
Occlusion Detection Time: Pressure limit at 2 PSI, Flow Rate at 1 mL/hr	8 minutes (2 PSI setting), 55:00 minutes (10 PSI setting)
Occlusion Detection Time: Pressure limit at 10 PSI, Flow Rate at 25 mL/hr	8 minutes (2 PSI setting), 108 seconds (1:48 minutes at 10 PSI setting), max pressure reached was 10.2 PSI
Occlusion Detection Time: Pressure limit at 10 PSI, Flow Rate at 100 mL/ hr	26 seconds (10 PSI setting), max pressure reached was 10.1 PSI
Occlusion Detection Time: Pressure limit at 10 PSI, Flow Rate at 500 mL/ hr	8 seconds (10 PSI setting), max pressure reached was 9.9 PSI
Occlusion Detection Time: Pressure limit at 10 PSI, Flow Rate at 999 mL/ hr	8 seconds (10 PSI setting), max pressure reached was 10.8 PSI
Post-Occlusion Pressure Alarm: Occlusion Relief Bolus Volume Delivered - Pressure limit at 2 PSI, Flow Rate at 25 mL/hr	Measured value was <0.7 mL after bolus release
Post-Occlusion Pressure Alarm: Occlusion Relief Bolus Volume Delivered - Pressure limit at 10 PSI, Flow Rate at 25 mL/hr	Measured value was 0.7 mL after bolus release
Battery Operating Time: Flow Rate at 25mL/hr (typical)	> 12 Hours at 25mL/hr Rate

AAMI ID 26 -1998 50.102 Data Set Flow Rate at T(0) 1 mL/hr



Figure G-1.



Figure G-2.







AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve

Figure G-4.

AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve over T(1) Period - 1 mL/hr



Figure G-5.





Figure G-6.

AAMI ID 26 -1998 50.102 Data Set Flow Rate at T(0) 25 mL/hr



Figure G-7.



Figure G-8.



AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve over T(1) Period - 25 mL/hr





AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve

Figure G-10.



AAMI ID 26 -1998 50.102 Data Set Flow Rate at T(0) 999 mL/hr

Figure G-11.

AAMI ID 26 -1998 50.102 Data Set Volume Delivery - 999 mL/hr 6500 6000 5500 5000 4500 (iii) 4000 3500 3000 2500 2000 1500 1000 500 0 0:00:00 0:11:00 0:22:00 0:33:00 0:44:00 0:44:00 1:06:00 1:17:00 1:28:00 1:39:00 1:50:00 2:01:00 2:45:00 2:55:00 3:07:00 3:45:00 3:45:00 3:40:00 4:46:00 4:57:00 5:08:00 5:19:00 5:30:00 5:31:00 5:41:00 5:52:00 6:03:00 6:03:00 2:12:00 2:23:00 2:34:00 3:51:00 4:02:00 4:13:00 4:24:00 4:35:00

G-8

Figure G-12.


AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve

Figure G-13.

AAMI ID 26 -1998 50.102 Data Set - Standard Trumpet Curve over T(2) Period - 999 mL/hr



Figure G-14.

APPENDIX H

1119 I.V. POLE ASSEMBLIES AND PARTS DESCRIPTIONS

General Precautions

- Federal law in the USA restricts this device to sale by or on the order of a physician.
- For safe operation, use only Iradimed Corporation recommended MRIcompatible or MRI-safe accessories.
- Always secure the IV Pole wheel locks after positioning within the MRI Magnet Room.
- Product damage may occur unless proper care is exercised during unpacking and installation. Use only recommended assemblies and parts. Any replacement component must be non-magnetic for safe operation.
- Refer all service to Iradimed Corporation, or Authorized Service Representatives.
- A maximum number of two (2) pumps can be safely used on any one (1) IV pole, mounted no more than 54 inches [137cm] from the floor.

NOTE: There are several 1119 I.V. Pole configurations. When using the following assembly instructions, please match the appropriate instructions to your specific 1119 I.V. Pole.

Routine Maintenance

Periodically check all mounting hardware. Tighten as necessary for optimal operation.

Cleaning the Mounting Assembly

1. The mounting assembly may be cleaned with most mild, non-abrasive solutions commonly used in the hospital environment (e.g. diluted bleach, ammonia, or alcohol solutions).

2. The surface finish will be permanently damaged by strong chemicals and solvents such as acetone and trichloroethylene.

3. Do not use steel wool or other abrasive material to clean the mounting assembly.

4. Damage caused by the use of unapproved substances or processes will not be covered by warranty. We recommended that you test any cleaning solution on a small area of the mounting assembly that is not visible to verify compatibility.

5. Never submerge the roll stand or allow liquids to enter the mounting assemblies Wipe any cleaning agents off the mounting assembly immediately, using a waterdampened cloth. Dry all mounting assemblies thoroughly after cleaning.

CAUTION: Iradimed Corporation makes no claims regarding the efficacy of the listed chemicals or processes as a means for controlling infection. Consult your hospital's infection control officer or epidemiologist. To clean or sterilize mounted instruments or accessory equipment, refer to the specific instructions delivered with those products.

Configuration 1 1119 I.V. POLE ASSEMBLIES AND PARTS DESCRIPTIONS



Installation Guide

Iradimed 1119 MRI Pump Roll Stand Kit

The purpose of this guide is to:

- 1. Describe assembly of Roll Stand Post and Base (page H-3).
- 2. Describe attachment of IV Bag Hooks to IV Pole (page H-3).
- 3. Describe assembly of IV Poles (page H-4).

General Precautions

- Federal law in the USA restricts this device to sale by or on the order of a physician.
- · For safe operation, use only Iradimed Corporation recommended MRI-compatible or MRI-safe accessories.
- Always secure the IV Pole wheel locks after positioning within the MRI Magnet Room.
- Product damage may occur unless proper care is exercised during unpacking and installation. Use only GCX Corporation recommended assemblies and parts. Any replacement component must be non-magnetic for safe operation.
- Refer all service to Iradimed Corporation, or GCX Corporation Authorized Service Representatives.
- A maximum number of two (2) pumps can be safely used on any one (1) IV pole, mounted no more than 54 inches [137cm] from the floor.

Parts Reference

The following parts and hardware are included in this installation kit (see illustrations for parts; hardware not shown):

ltem #	Description	Qty
1	IV Pole, 66" 2-piece	1
2	Base (with 10 lb. Counterweight)	1
3	5/16-18 x 1" Hex Head Cap Screw (HHCS)	1
4	5/16 Flat Washer	1
5	5/16 Split Lock Washer	1
6	Hook, IV Bag	2
7	1/4-20 x 3/4" Flat Head Socket Cap Screw (FHSCS)	1
8	Tree Disc	1
9	5/32" Hex Wrench	1
10	3/32" Hex Wrench	1



Tools Required

1/2" [13mm] wrench (not provided) 5/32" and 3/32" hex wrenches (provided).

1119 Assembly Instruction Rev. C 010308

Assembling the Roll Stand (Post to Base)

- 1. Insert Lower IV Pole in Base and lay assembly on its side for access to bottom of Base (not shown).
- 2. Using a 1/2" [13 mm] wrench, fasten Post to Base with one (1) 5/16-18 x 2" HHCS, 5/16 split lock washer, and 5/16" flat washer.



Attaching IV Bag Hooks to Upper IV Pole

- 1. Assemble Tree Disc over IV Bag Hooks. Installation Note: If IV Bag Hooks do not fit easily into Tree Disc slots, gently tap Hooks into slots with a rubber mallet.
- Align locator pin (top of Pole) with slot in Tree Disc. Using the 5/32" hex wrench provided, fasten assembly to top of Upper IV Pole with one (1) 1/4-20 x 3/4" FHSCS (1) as shown below.



Attaching Upper IV Pole to Lower Pole

- 1. Screw Upper Pole clockwise (CW) onto Lower Pole. Tighten as much as possible by hand.
- 2. Using the 3/32" hex wrench provided, tighten the set screw in the Upper Pole.



Routine Maintenance

Periodically check all mounting hardware. Tighten as necessary for optimal operation.

Cleaning the Mounting Assembly

- 1. The mounting assembly may be cleaned with most mild, non-abrasive solutions commonly used in the hospital environment (e.g. diluted bleach, ammonia, or alcohol solutions).
- 2. The surface finish will be permanently damaged by strong chemicals and solvents such as acetone and trichloroethylene.
- 3. Do not use steel wool or other abrasive material to clean the mounting assembly.
- 4. Damage caused by the use of unapproved substances or processes will not be covered by warranty. We recommended that you test any cleaning solution on a small area of the mounting assembly that is not visible to verify compatibility.
- 5. Never submerge the roll stand or allow liquids to enter the mounting assemblies Wipe any cleaning agents off the mounting assembly immediately, using a water-dampened cloth. Dry all mounting assemblies thoroughly after cleaning.

CAUTION: GCX makes no claims regarding the efficacy of the listed chemicals or processes as a means for controlling infection. Consult your hospital's infection control officer or epidemiologist. To clean or sterilize mounted instruments or accessory equipment, refer to the specific instructions delivered with those products.

Configuration 2 1119 I.V. POLE ASSEMBLIES AND PARTS DESCRIPTIONS

NOTE: The following tools are required for the assembly of this IV Pole:

Small Screws1/8" HEXBig Screws3/16" HEXBolthead1/2" Wrench







NOTES